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(71) Applicant (for all designated States except US): CHIRON DIAGNOSTICS CORPORATION [US/US]; 333 Coney Street, East Walpole, MA 02032 (US).

(72) Inventors; and

- (75) Inventors/Applicants (for US only): SANDHU, Gurpreet, S. [IN/US]; Apartment 212, 10320 Devonshire Circle, Bloomington, MN 55431 (US). KLINE, Bruce, C. [US/US]; 2315 Kline Lane, S.W., Rochester, MN 55902 (US).
- (74) Agents: MORGENSTERN, Arthur, S.; Chiron Diagnostics Corporation, 63 North Street, Medfield, MA 02052 (US) et al.

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(54) Title: NUCLEIC ACID PROBES FOR THE DETECTION AND IDENTIFICATION OF FUNGI

(57) Abstract

Nucleic acid probes and primers are described for detecting fungi that cause disease in humans and animals, as well as spoilage of food and beverages. These probes can detect rRNA, rDNA or polymerase chain reaction products from a majority of fungi in clinical, environmental or food samples. Nucleic acid hybridization assay probes specific for *Paracoccidioides brasiliensis* and *Pneumocystis carinii* are also described.

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NUCLEIC ACID PROBES FOR THE DETECTION AND IDENTIFICATION OF FUNGI

FIELD OF INVENTION

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The inventions described and claimed herein relate to the design and composition of two nucleic acid probes capable of detecting many different fungal organisms in clinical, food, environmental and other samples. The inventions described and claimed herein also relate to the design and composition of probes capable of specifically detecting and identifying Acremonium sp., Aspergillus clavatus, Aspergillus flavus, Aspergillus fumigatus, Aspergillus glaucus, Aspergillus nidulans, Aspergillus niger, Aspergillus ochraceus, Aspergillus terreus, Aspergillus unguis, Aspergillus ustus, Beauveria sp., Bipolaris sp., Blastoschizomyces sp., Blastomyces dermatitidis, Candida albicans, Candida glabrata, Candida guilliermondii, Candida kefyr, Candida krusei, Candida lusitaniae, Candida parapsilosis, Candida tropicalis, Chrysosporium sp., Cladosporium sp., Coccidioides immitis, Cryptococcus neoformans var gattii serotype B, Cryptococcus neoformans serotype A, Cryptococcus laurentii, Cryptococcus terreus, Curvularia sp., Fusarium sp., Filobasidium capsuligenum, Filobasidiella (Cryptococcus) neoformans var bacillispora serotype C, Filobasidiella (Cryptococcus) neoformans var neoformans serotype D, Filobasidium uniguttulatum, Geotrichum sp., Histoplasma capsulatum, Malbranchea sp., Mucor sp., Paecilomyces sp., Paracoccidioides brasiliensis, Penicillium species, Pneumocystis carinii, Pseudallescheria boydii, Rhizopus sp., Sporothrix schenkii, Scopulariopsis brevicaulis sp., Scopulariopsis brumpti, Saccharomyces cerevisiae, and Trichosporon beigelii in clinical, food, environmental and other samples.

Fungi are eukaryotic microorganisms that are universally distributed. While in nature fungi play a major role in the decomposition of plant materials, they are also responsible for spoilage of food, beverage and pharmaceutical preparations. Out of an

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estimated 100,000 species of fungi described by mycologists, approximately 150 species are pathogenic to man and animals. The increasing incidence of AIDS and the development of newer treatments for hematologic malignancies and organ transplants has lead to an increase in the number of immunocompromised patients. These patients have a high risk of developing fungal infections, which if not rapidly diagnosed and treated are capable of causing death in a matter of days. The number of antifungal drugs is limited and their toxic side effects on the patient are much higher than that of comparable antibacterial therapy. A rapid diagnosis of fungal infection and start of treatment is critical in these patients. Books by Kwon-Chung and Bennett, along with Sarosi and Davies, provide an overview into the medical importance of fungi.

Fungal organisms are identified by morphology and nutritional characteristics. Fungi may take anywhere from two days to several weeks to grow in culture and often the same organism can take radically different forms depending on the growth conditions. This makes timely identification difficult even for the classically trained expert and impedes the treatment of patients where rapid identification of genus and species is of medical advantage.

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The incidence and distribution of major pathogenic fungi varies by geographic location. Aspergillus fumigatus, Blastomyces dermatitidis, Candida albicans, Coccidiodes immitis, Cryptococcus neoformans, Histoplasma capsulatum, Paracoccidioides brasiliensis, Pseudallescheria boydii and Sporothrix schenkii represent some of the leading causes of mycotic infections.

Aspergillus fumigatus is among the top three causes of systemic fungal infection treated in hospitals. It usually affects patients with organ transplants, acute leukemias and burns and can be rapidly fatal if not diagnosed quickly. With over 150 species of Aspergillus present in the soil, air and water, accurate detection of Aspergillus fumigatus becomes extremely important. Aspergillus clavatus, Aspergillus flavus, Aspergillus fumigatus, Aspergillus glaucus, Aspergillus nidulans, Aspergillus niger, Aspergillus ochraceus, Aspergillus terreus, Aspergillus unguis and Aspergillus ustus represent a majority of Aspergillus species seen in clinical specimens and their presence can cause diagnostic difficulties. Aspergillus flavus, Aspergillus fumigatus and Aspergillus niger have been linked with disease in humans, with Aspergillus fumigatus being the predominant pathogen in North America. A few immunologic

tests exist for Aspergillus fumigatus but these have limited sensitivity and specificity. There are also reports of development of polymerase chain reaction based tests for Aspergillus fumigatus based on the amplification of the Asp f1 antigen gene and a ribosomal intergenic spacer (Spreadbury et. al.). The Spreadbury technique is based on the PCR amplification of a 401 bp fragment spanning the large subunit rRNA/intergenic spacer region. This relies on a pair of primers to specifically amplify DNA from Aspergillus fumigatus only, and is of no utility in identifying other fungi.

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Blastomyces dermatitidis is present in the soil, usually in bird droppings and animal feces. Infections often occur at construction sites and the ensuing lung infiltration and pneumonitis are usually fatal in immunocompromised patients. Diagnosis by culture may take weeks, and the organism is occasionally mistaken for other fungi. Existing immunological diagnostic tests are unreliable, and there is a need for rapid and reliable DNA based diagnostic tests. Similarly, Histoplasma capsulatum exists in the soil and is known to have infected at least 20% of the population of North America. Most infections start in the lung and resolve spontaneously, but may occasionally spread to other organs. AIDS patients represent a growing number of cases of Histoplasmosis. Diagnosis is difficult as immunological tests are often negative during the first 4-6 weeks of infection. Coccidioides immitis is found in abundance in the soil in Southwestern United States. Dust storms, farming, building construction, earthquakes and even hiking have been linked with outbreaks of disease. Lung infection followed by cavitation and disseminated miliary coccidioidomycosis are seen. Meningitis is usually lethal, and as with other fungi, mortality is highest in debilitated hosts. Four serotypes of Cryptococcus neoformans cause disease in humans. These are Cryptococcus neoformans serotype A, Cryptococcus neoformans var gatti serotype B, Filobasidiella (Cryptococcus) neoformans var bacillispora serotype C and Filobasidiella (Cryptococcus) neoformans var. neoformans serotype D. The incidence of this disease is growing rapidly, with up to 10% of HIV infected people developing cryptococcosis. DNA probes capable of detecting all 4 serotypes are required for the early diagnosis and treatment for life threatening infections like cryptococcal meningitis. A report by Stockman et. al. discusses commercial tests for Histoplasma, Blastomyces, Coccidioides, and Cryptococcus based on the 18S rRNA (Gen-Probe, Inc., San Diego, CA). The authors report sensitivities ranging from 87.8

to 100% and a specificity of 100%. One drawback of these probes is that these are used on rRNA extracted from fungal cultures. As some fungi may require up to 3 weeks to grow in culture, this technique cannot be used to expedite diagnosis until a culture becomes available.

Candida albicans is one of the most common causes of fungal infection in humans. It is present in the respiratory, gastrointestinal and female genital tract of healthy individuals, and acts as an opportunistic pathogen in debilitated individuals on steroid or chemotherapy. Diabetes mellitus and indwelling catheters are other predisposing causes. Immunocompromised hosts show rapid hematogenous spread of fungi. Morbidity and mortality in untreated cases is high. Candida glabrata, Candida guilliermondii, Candida kefyr, Candida krusei, Candida lusitaniae, Candida parapsilosis and Candida tropicalis are also known to cause disease in humans. DNA probes capable of identifying these individual species would eliminate the need for multiple blood cultures and lengthy biochemical speciation.

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Paracoccidioidomycosis, which is a deep-seated systemic infection of humans, is a major health problem in Central and South America. The disease is caused by *Paracoccidioides brasiliensis*, a thermally dimorphic fungus. Classically, diagnosis has been made by detecting yeast cells with multiple buds, the distinctive "pilots wheel" morphology. Unfortunately, such forms are not always observed in clinical materials. Histologically, the yeast form of this species is sometimes difficult to distinguish from that of *Histoplasma capsulatum*, *Blastomyces dermatitidis*, *Coccidioides immitis*, and *P. loboi*. While *P. brasilienis* produces a glycoprotein antigen than is also diagnostic, the antigen is also produced by *P. loboi*. *P. brasilienis* can be cultured in vitro; however *P. loboi* cannot be cultured. *P. brasiliensis* is a slow growing organism that can take one to three weeks to culture for identification. Clearly, the existence of a species specific probe against *P. brasiliensis* would be of significant value for rapid detection and identification.

Pneumocystis carinii is a common and major opportunistic pathogenic fungus that infects immunocompromised patients. More than 80% of HIV infected patients will develop *P. carinii* pneumonia if not prophylactically treated to prevent infection. Therefore, this fungus is widely recognized as one of the most important causes of morbidity and mortality in AIDS patients. To date, polymerase chain reaction primers

directed at six different gene targets have been developed. These include 5S and 18 rDNA genes, an intragenic transcribed spacer sequence between ribosomal genes, thymidylate synthase, dihydrofolate reductase, and large ribosomal rDNA gene of mitochondria. However, each set of primers requires a unique and specific set of PCR conditions to function optimally (Lu et al., J. Clin. Microbial. 33:2785-2788, 1995).

Recent advances in molecular techniques have led to the approach of microbe detection and identification based upon the DNA sequence of ribosomal genes. Commonly used detection techniques include either direct amplification of the ribosomal DNA (rDNA) genes by the polymerase chain reaction, or reverse transcription of the ribosomal RNA (rRNA) into complementary DNA (cDNA) followed by polymerase chain reaction amplification of the cDNA. Ribosomes are composites of unique rRNA and protein species that function in the translation of messenger RNA into protein. Evolutionary studies are consistent with the interpretation that all extant life has evolved from a single organism. Thus, all cellular organisms contain rRNA and these rRNAs are related by evolution. The evolutionary process is such that each species of organism appears to have unique regions of sequence in its ribosomal genes. The presence of these unique species specific regions allows one to design DNA probes that under conditions of hybridization will specifically bind to, and identify the polymerase chain reaction amplified DNA from only one species of fungus. For the purposes of this application, the word "primer" is used to mean a nucleotide sequence which can be extended by template-directed polymerization, and "probe" is used to mean a nucleotide sequence capable of detecting its complementary sequence by hybridization. Also, for the purpose of this application, the phrase "nucleotide sequence" is intended to include either DNA or RNA forms or modification thereof. Furthermore, those versed in the art will recognize that primer sequences can be used as probes and vice versa. The use of nucleic acid hybridization to detect specific nucleic acid sequences of interest is also described by Kohne (U.S. Patent 4,851,330, 7/1989).

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In prokaryotes and eukaryotes, ribosomal RNA and the corresponding rDNA genes are identified by the size of the RNA. The sizes are related in terms of sedimentation velocity or S values. Thus, for prokaryotes the values are 5S, 16S, and 23S; and for eukaryotes the values are 5S, 5.8S, 18S and 28S. Because all ribosomes

perform the same function which is essential for cell viability, ribosomal sequences are largely conserved, yet certain regions of each ribosomal species are subject to more variation without consequence to function. It is these hypervariable regions that allow one to identify different species amongst members of the same genus. As noted in the references, there are several reports where 5S, 18S and the intergenic spacer between 5.8S and 28S rDNA have been used for the detection and identification of fungi (Holmes et. al., Hopfer et. al., Lott et. al., Maiwald et. al., Makimura et. al., Mitchell et. al., Nakamura et. al.). Holmes et. al. describe a PCR test based on the coamplification of the 5S rDNA and an adjacent nontranscribed spacer region. This identifies only Candida albicans and detects other Candida species without identifying individual organisms. Hopfer et. al. and Maiwald et. al. both use universal primers to amplify 18S rDNA from several fungi including Candida sp., Aspergillus fumigatus, Cryptococcus neoformans and Trichosporon sp. These amplicons are digested with restriction enzymes and the cut fragments are sized by gel electrophoresis. This restriction fragment length polymorphism pattern enables them to identify most but not all organisms. This technique can be used on amplified DNA from a pure fungal culture. As clinical samples such as sputum usually contain multiple fungal organisms, this technique has little utility in diagnosis as multiple overlapping fragments obtained from a mix of fungi would be nearly impossible to interpret. Lott et. al. use the 5.8S RNA and the internal transcribed spacer (ITS2) to identify and speciate Candida albicans and related Candida species. Makimura amplifies a 687 bp fragment from the 18S rDNA of 25 medically important fungi and uses these in the diagnosis of Candida albicans in clinical samples. Mitchell uses nested PCR to amplify 5.8S and internal transcribed spacer (ITS) to identify Cryptococcus neoformans. No subsequent testing is done to verify the identity of the amplified DNA. Nakamura et. al. use 18S primers to detect Aspergillus fumigatus infections of the lung. Most protocols given in these references can only be used to detect an extremely limited number of fungi from a clinical specimen. Hopfer et. al. and Maiwald et. al. can identify multiple organisms from pure cultures, but their utility for clinical specimens containing multiple fungal species is limited at best.

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United States patents have been issued to Weisburg et. al. for probes developed for the detection of 18S small subunit ribosomal RNA sequences in fungi.

These probes will detect fungi from many species, but cannot be used easily to identify any single species. United States patents have also been issued to Milliman for probes developed for the specific detection of the bacteria Staphylococcus aureus based on the 16S ribosomal sequences. Hogan et. al. (European Pat. App. 0,272,009) describe one fungal probe for 18S rRNA and three fungal probes for 28S rRNA sequences. Two of these 28S probes detect several different fungi while the third probe detects Candida krusei from a limited panel of 10 fungi. None of the 28S probes described by Hogan et. al. is related to any of the probes described in our invention. All probes claimed in our invention can be mapped within the first 900 base pairs of a 28S gene. The probes described by Hogan et. al. are located further 3' on the 28S sequence, between base pairs 1000 and 2000 (these numbers are comparable to the primary sequence of Saccharomyces cerevisiae 28S rRNA gene. Genbank accession number: J01355). Leclerc et. al. have published reports analyzing the phylogenetic relationship between fungi based on partial DNA sequences of several fungal 28S genes sequenced by them. Some of the organisms claimed to have been sequenced by Leclerc are the same as some organisms sequenced by us. These are Sporothrix schenckii, Pseudallescheria boydii, Blastomyces dermatitidis, Histoplasma capsulatum and Chrysosporium sp. Leclerc et. al. have not published any sequence data in their report, and to the best of our knowledge, they have not made these sequences publically available. The reverse-complement sequence of their sequencing primer 401 (TCCCTTTCAA CAATTTCACG) overlaps our SEQ ID NO: 1 (GTGAAATTGT TGAAAGGGAA) by 19 nucleotides and their sequencing primer 636 (GGTCCGTGTT TCAAGACGG) overlaps our SEQ ID NO: 2 (GACTCCTTGG TCCGTGTT) by 10 nucleotides. We are aware of no reports in the literature of variable regions from 28S rRNA genes of fungi being used as targets for the development of species specific diagnostic probes.

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As discussed above, most present techniques for the molecular detection of fungi rely on the use of highly specific primers for the PCR amplification of only one fungal species. Those that employ "Universal" primers for a PCR amplification of DNA from multiple organisms, use post-PCR amplicon identification techniques that are useful only on pure cultures of fungi. These are not be able to identify fungi from a clinical specimen containing multiple fungal organisms. Our first aim was to

develop "Universal" primers for the 28S gene. These primers would be capable of amplifying in a PCR, 28S rDNA from most fungi. Our subsequent aim was to develop species specific probes for fungi of interest, that would be used to analyze our "Universal" 28S amplicon. These species specific probes would be able to detect the presence of fungi of interest even in situations containing mixed fungal species.

One aspect of this invention is to provide nucleic acid primers capable of detecting 28S sequences from DNA or RNA of most fungi. These would be used as "Universal" primers in a polymerase chain reaction to amplify 28S sequences from any fungus present in clinical, food, environmental or other samples. These "Universal" primers would also be used to sequence the amplified DNA. The sequence obtained would be used to identify the fungus by comparing with a database of known fungal sequences.

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A second aspect of this invention is to provide nucleic acid probes capable of detecting and identifying, by nucleic acid hybridization, the pathogens Aspergillus fumigatus, Blastomyces dermatitidis, Candida albicans, Coccidioides immitis, Cryptococcus neoformans, Histoplasma capsulatum, Aspergillus flavus, Aspergillus glaucus, Aspergillus niger, Aspergillus terreus, Candida glabrata, Candida guilliermondii, Candida kefyr, Candida krusei, Candida lusitaniae, Candida parapsilosis, Candida tropicalis, Paracoccidioides brasiliensis, Pneumocystis carinii, Pseudallescheria boydii, Sporothrix schenckii and other species by use of any of several different formats. Additionally, nucleotide sequence information is provided to identify these pathogens and other fungi by DNA sequence comparison (Table 3) or by the construction of additional probes.

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SUMMARY OF THE INVENTION

Nucleic acid probes and primers are described for detecting fungi that cause disease in humans and animals, as well as spoilage of food and beverages. These probes can detect rRNA, rDNA or polymerase chain reaction products from a majority of fungi in clinical, environmental or food samples. Nucleic acid hybridization assay probes specific for Aspergillus fumigatus, Blastomyces dermatitidis, Candida albicans, Coccidioides immitis, Cryptococcus neoformans, Histoplasma capsulatum, Aspergillus flavus, Aspergillus glaucus, Aspergillus niger, Aspergillus terreus, Candida glabrata, Candida guilliermondii, Candida kefyr, Candida krusei, Candida lusitaniae, Candida parapsilosis, Candida tropicalis, Paracoccidioides brasiliensis, Pneumocystis carinii, Pseudallescheria boydii, Sporothrix schenckii and other species (Table 1 and Table 3) are also described.

Figure 1 represents the relative position of the sequences described on the 28S subunit of fungi.

DETAILS OF THE INVENTION

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Our first objective was to develop nucleic acid primers for use in a polymerase chain reaction to amplify 28S genes from all fungi likely to be present in a clinical sample. This amplified DNA would then be amenable to probing with several different species specific probes. Each one of these species specific probes would, under conditions of hybridization, anneal to 28S ribosomal DNA from only one species of fungus, thereby detecting and identifying the species of fungus present in the clinical sample. The 28S gene was selected as a target because it had regions that were conserved among fungi and these would provide potential annealing sites for "universal" fungal probes. The ribosomal 28S genes were also expected to have hypervariable regions that would be unique enough to provide sites for species specific probes. The large rRNA gene is called the 23S rRNA gene in prokaryotes and 28S in eukaryotes. This designation is based on the length and therefore the sedimentation coefficient of these rRNA molecules. Fungal large subunit rRNAs vary in size among different organisms and are often referred to as being 25S, 26S or 28S.

Since fungi are eukaryotes, and to maintain uniformity in this application, we shall refer to fungal large subunit rRNA as 28S rRNA.

Published sequences from Cryptococcus neoformans, two Candida albicans, Saccharomyces cerevisiae and two Schizosaccharomyces pombe 28S genes are approximately 3.5 kilobases in length (Genbank accession numbers: L14068, L28817, X70659, J01355, Z19136 & Z19578). These four sequences were aligned, and a region of sequence variability was found clustered between coordinates 200 and 700 from the 5' end of these genes. As an initial starting point, two nucleic acid primers P1 (ATCAATAAGC GGAGGAAAAG) (SEQ ID NO:79) and P2 (CTCTGGCTTC ACCCTATTC) (SEQ ID NO:80) (see figure 1), capable of hybridizing to all 4 of the above mentioned organisms and not to human 28S sequences (GenBank accession number: M11167), were designed and used under low stringency hybridization conditions in a polymerase chain reaction to amplify approximately 800 base pairs of DNA spanning this hypervariable region from the following 34 fungi that were obtained from the Mayo Clinic fungal collection: Acremonium sp., Aspergillus 15 clavatus, Aspergillus fumigatus, Aspergillus glaucus, Aspergillus nidulans, Aspergillus niger, Aspergillus ochraceus, Aspergillus terreus, Aspergillus unguis, Aspergillus ustus, Beauvaria sp., Bipolaris sp., Blastomyces dermatitidis, Candida albicans, Candida glabrata, Candida guilliermondii, Candida kefyr, Candida krusei, Candida lusitaniae, Candida parapsilosis, Candida tropicalis, Chrysosporium sp., 20 Cladosporium sp., Coccidioides immitis, Cryptococcus neoformans serotype A, Curvularia sp., Geotrichum sp., Histoplasma capsulatum, Mucor sp., Penicillium sp., Pseudallescheria boydii, Saccharomyces cerevisiae, Sporothrix schenkii and Trichosporon beigelii.

DNA was extracted from the fungi listed above by the following method. A loopful of fungal culture was scraped off a culture plate using a sterile inoculation loop. The fungus was added one milliliter of sterile water in a 1.5 ml Sarsted (Newton, North Carolina) screw cap microcentrifuge tube. This tube was placed in a boiling water bath for 20 minutes in order to lyse the fungus and release DNA from the cells.

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Two microliters of this whole cell lysate was used in a PCR to amplify 28S rDNA. All PCR amplifications were carried out as hot-start reactions in a 50 ul reaction volume using Perkin-Elmer (Norwalk, CT) 0.5 ml thin-wall polypropylene tubes and a Perkin-

Elmer thermal cycler. Reagents added to the tube initially were 2.5 ul of 10X PCR buffer (100 mM tris pH 8.3, 500 mM KCl, 15 mM MgCl₂), 5.0 ul of 50% glycerol/1 mM cresol red, 8.0 ul of dNTP mix (1.25 mM each of dATP, dGTP, dTTP and dCTP), 12 picomoles of each nucleic acid primer and sterile water to make up a volume of 25 ul. A wax bead (Ampliwax Gem-100, Perkin-Elmer) was added and the tubes heated to 77°C for 1 minute and cooled to room temperature to form a wax barrier. 2.5 ul of 10X PCR buffer, 5.0 ul of 50% glycerol/1 mM cresol red, 0.2 ul Taq polymerase (AmpliTaq 5U/ul, Perkin-Elmer) and 15.3 ul of sterile water was added to the tube along with 2.0 ul of DNA from the fungal whole cell lysate described above. 50 cycles of thermal cycling was carried out at 94°C - 30 sec, 40°C - 1 min, 72°C - 2 min. The amplified DNA was electrophoresed and purified from a low melt agarose gel by tris buffered phenol pH 8.0, phenol/chloroform/isoamyl alcohol (25:24:1 by vol.) and 3 ether extractions, followed by isopropanol precipitation and 70% ethanol wash.

We completely sequenced both strands of DNA amplified from the organisms listed above. All sequencing was carried out on an Applied Biosystems 373A sequencer. Every nucleotide in the sequences generated was verified and confirmed by examining the complementary nucleotide from the second strand sequence. We had now created a novel database consisting of nucleic acid sequences spanning a variable region of the 28S rDNA from a diverse collection of medically important fungi.

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While the complete sequences for Candida albicans, Cryptococcus neoformans and Saccharomyces cerevisiae 28S genes had previously been published and deposited in GenBank, it was not obvious, nor had it been defined, whether any regions of sequence identity among these three organisms would also be conserved among all fungi of interest. DNA sequences from all the fungi in our novel 28S database had to be analyzed in order to develop "Universal" 28S probes. All sequences were subjected to extensive manipulation to identify optimal relative allignments in order to identify regions of similarity for use as "Universal" probes. The selected probe sequences had to meet several important criteria besides the condition of being present in 28S genes from most fungal species. Each probe sequence required an appropriate thermal profile, secondary structure and utility in a

DNA amplification reaction. These probes were optimized to work for PCR amplification in pure cultures of fungus, as well as in the presence of DNA from multiple sources as in the case of clinical specimens. The probes were also designed to facilitate direct sequencing of the amplified DNA. Our analysis led to the discovery of the oligonucleotide probes listed in (SEQ ID NO:1) and (SEQ ID NO:2). (For their location, see Figure 1.) The successful identification of these two probes ((SEQ ID NO:1) and (SEQ ID NO:2)) completed our first objective to develop nucleic acid probes that would hybridize to, and detect 28S rRNA and rDNA from a majority of fungi (Figure 1 and Table 1). As shown later in this application, the novel sequence information generated by the use of our "Universal" probes allowed us to develop species-specific probes ((SEQ ID NO:3) to (SEQ ID NO:23), (SEQ ID NO:75) and (SEQ ID NO:76)) capable of identifying 21 different disease-causing fungi.

Table 1:

Presence of hybridization sites for probes SEQ ID NO: 1 and SEQ ID NO: 2 in 28S nucleic acid sequences.

	SEQ ID NO: 1	SEQ ID NO: 2
Acremonium sp.	+	+
Aspergillus clavatus	+	+
Aspergillus flavus	+	+
Aspergillus fumigatus	+	+
Aspergillus glaucus	+	+
Aspergillus nidulans	. +	+
Aspergillus niger	+	+
Aspergillus ochraceus	+ .	+
Aspergillus terreus	+	+
Aspergillus unguis	+	+
Aspergillus ustus	+	+
Beauvaria sp.	+	+

Bipolaris sp.	+	+
Blastomyces dermatitidis	+	+
Blastoschizomyces sp.	+	+
Candida albicans	+	+
Candida glabrata	+	+
Candida guilliermondii	+	+
Candida kefyr	+	+
Candida krusei	+	+
Candida lusitaniae	+	+
Candida parapsilosis	+	+
Candida tropicalis	+	-+
Chrysosporium sp.	+	+
Cladosporium sp.	+	+
Coccidioides immitis	+	+
Cryptococcus laurentii	+	+
Cryptococcus neoformans serotype	+	+
A		
Cryptococcus neoformans var.	+	+
gattii serotype B	•	
Cryptococcus terreus	+	+ .
Curvularia sp.	+	+
Filobasidiella (Cryptococcus)	+	+
neoformans var bacillispora		
serotype C		
Filobasidiella (Cryptococcus)	· +	+
neoformans var neoformans		
serotype D		
Filobasidium capsuligenum	+	+
Filobasidium uniguttulatum	+	+
Fusarium sp.	+	+
Geotrichum sp.	+	+
	<u> </u>	L

Histoplasma capsulatum	+	+ .
Malbranchea sp.	+	+
Mucor sp.	+	+
Paecilomyces sp.	+	+
Penicillium sp.	+	+
Pseudallescheria boydii	+	+ .
Rhizopus sp.	+	+
Saccharomyces cerevisiae	+	+
Scopulariopsis brevicaulis	+	+
Scopulariopsis brumptii	+	+
Sporothrix schenckii	+	+
Trichosporon beigelii	+	+
Paracoccidioides brasiliensis	+	+
Pneumocystis carinii	+	+
Human	-	+

Probes SEQ ID NO: 1 and SEQ ID NO: 2 were used to successfully amplify (Table 2) and sequence DNA (Table 3) spanning this variable region from the following 51 organisms: Acremonium sp., Aspergillus clavatus, Aspergillus flavus, Aspergillus fumigatus, Aspergillus glaucus, Aspergillus nidulans, Aspergillus niger, Aspergillus ochraceus, Aspergillus terreus, Aspergillus unguis, Aspergillus ustus, Beauvaria sp., Bipolaris sp., Blastomyces dermatitidis, Blastoschizomyces sp., Candida albicans, Candida glabrata, Candida guilliermondii, Candida kefyr, Candida krusei, Candida lusitaniae, Candida parapsilosis, Candida tropicalis, Chrysosporium sp., Cladosporium sp., Coccidioides immitis, Cryptococcus neoformans serotype A, Cryptococcus neoformans var. gattii serotype B, Cryptococcus terreus, Cryptococcus laurentii, Curvularia sp., Filobasidiella (Cryptococcus) neoformans var bacillispora serotype C, Filobasidiella (Cryptococcus) neoformans var neoformans serotype D, Filobasidium capsuligenum, Filobasidium uniguttulatum, Fusarium sp., Geotrichum sp., Histoplasma capsulatum, 15 Malbranchea sp., Mucor sp., Paecilomyces sp., Penicillium sp., Pseudallescheria

boydii, Rhizopus sp., Saccharomyces cerevisiae, Scopulariopsis brevicaulis, Scopulariopsis brumptii, Sporothrix schenkii, Paracoccidioides brasiliensis, Pneumocystis carinii, and Trichosporon beigelii. This list contains all 4 serotypes (A, B, C and D) of Cryptococcus neoformans. This sequence information generated by the use of probes SEQ ID NO: 1 and SEQ ID NO: 2 expanded the size of our database consisting of fungal 28S sequences. All amplified DNA was sequenced across both strands from a minimum of two different isolates of each organism to ensure accuracy of the data generated.

Table 2:
Polymerase chain reaction amplification of 28S rDNA with probes SEQ ID NO: 1 and SEQ ID NO: 2.

	PCR with SEQ ID
	NO: 1 & NO: 2
Acremonium sp.	+
Aspergillus clavatus	+
Aspergillus flavus	+
Aspergillus fumigatus	+
Aspergillus glaucus	+
Aspergillus nidulans	+
Aspergillus niger	+
Aspergillus ochraceus	+
Aspergillus terreus	+
Aspergillus unguis	+
Aspergillus ustus	+
Beauvaria sp.	+
Bipolaris sp.	+
Blastomyces dermatitidis	+
Blastoschizomyces sp.	+
Candida albicans	+

Candida glabrata	+
Candida guilliermondii	+
Candida kefyr	+
Candida krusei	+
Candida lusitaniae	+
Candida parapsilosis	+
Candida tropicalis	+
Chrysosporium sp.	+
Cladosporium sp.	+
Coccidioides immitis	+
Cryptococcus laurentii	+
Cryptococcus neoformans serotype	+
A	
Cryptococcus neoformans var.	+
gattii serotype B	
Cryptococcus terreus	+
Curvularia sp.	+
Filobasidiella (Cryptococcus)	+
neoformans var bacillispora	
serotype C	
Filobasidiella (Cryptococcus)	. +
neoformans var neoformans	
serotype D	
Filobasidium capsuligenum	+
Filobasidium uniguttulatum	+
Fusarium sp.	+
Geotrichum sp.	+
Histoplasma capsulatum	+
Malbranchea sp	+
Mucor sp.	+
Paecilomyces sp.	+
	

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Penicillium sp.	+
Pseudallescheria boydii	+
Rhizopus sp.	+.
Saccharomyces cerevisiae	+
Scopulariopsis brevicaulis	+
Scopulariopsis brumptii	+
Sporothrix schenckii	+
Trichosporon beigelii	+
Paracoccidioides brasiliensis	+
Pneumocystis carinii	+
Human	-

This list of fungi sequenced by us represents organisms responsible for most cases of subcutaneous and deep mycotic infections in humans and also includes saprophytes (non-pathogenic fungi) commonly encountered in clinical isolates. Since the two probes (SEQ ID NO: 1 and SEQ ID NO: 2) hybridize to 28S rDNA from all the fungi listed above, they are capable of diagnosing the presence of a majority of fungi that are likely to be present in a clinical specimen. They are believed to be primers for universally detecting fungi.

Probes listed in SEQ ID NO: 1 and SEQ ID NO: 2 were also checked for their potential ability to hybridize to, and amplify (in a polymerase chain reaction) 23S sequences from bacteria by searching for hybridization sites among the 539 bacterial 23S genes listed in GenBank. Bacterial 23S rDNAs do not have suitable hybridization sites for SEQ ID NO: 1 and SEQ ID NO: 2 and these two probes should not be able to amplify bacterial DNA under stringent conditions.

Our second objective was to develop species specific probes, which under hybridization conditions, would detect Aspergillus fumigatus, Blastomyces dermatitidis, Candida albicans, Coccidioides immitis, Cryptococcus neoformans, Histoplasma capsulatum, Aspergillus flavus, Aspergillus glaucus, Aspergillus niger, Aspergillus terreus, Candida glabrata, Candida guilliermondii, Candida kefyr, Candida krusei, Candida lusitaniae, Candida parapsilosis, Candida tropicalis,

Paracoccidioides brasiliensis, Pneumocystis carinii, Pseudallescheria boydii, and Sporothrix schenckii. We used our database of fungal 28S nucleic acid sequences to create a multiple sequence alignment of all the organisms that we had sequenced. Every individual sequence was subjected to intensive comparison with all other sequences in our database in order to discover unique regions of sequence that would be present only in the fungus of interest, and would be absent in all other fungi. When unique stretches of sequence were identified, these were further analyzed for thermal profile and secondary structure. Each probe constructed by us will, under conditions of hybridization, specifically hybridize to and detect, nucleic acid sequence from the unique region of only one specific target fungus. Those versed in the art will recognize that specification of a single-stranded DNA sequence implies the utility of the complementary DNA sequence, as well as the two equivalent RNA sequences. Furthermore, sequences incorporating modification of any of the mojeties comprising the nucleic acid (i.e., the base, the sugar or the backbone) are functional equivalents of the sequence. It should also be recognized that these additional sequences can potentially serve as probes or primers. Finally, those versed in the art recognize that comparisons of extensive DNA sequences provides enough variability and uniqueness to speciate organisms (Table 3).

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The nucleic acid sequences for these species specific synthetic probes are listed in SEQ ID NO: 3 to SEQ ID NO: 23, SEQ ID NO:75 and SEQ ID NO:76.

There are two probes specific for *Cryptococcus neoformans*, two probes specific for *Sporothrix schenckii*, and one probe each for *Aspergillus fumigatus*, *Blastomyces dermatitidis*, *Candida albicans*, *Coccidioides immitis*, *Histoplasma capsulatum*, *Aspergillus flavus*, *Aspergillus glaucus*, *Aspergillus niger*, *Aspergillus terreus*, *Candida glabrata*, *Candida guilliermondii*, *Candida kefyr*, *Candida krusei*, *Candida lusitaniae*, *Candida parapsilosis*, *Candida tropicalis*, *Paracoccidioides brasiliensis*, *Pneumocystis carinii*, and *Pseudallescheria boydii* 28S rRNA and rDNA. (See Tables 4 - 8 and further discussion below.)

All species specific probes developed by us are novel and to the best of our knowledge have not been reported in the literature. While all 28S genes sequenced by us had several regions that were different among the various species analyzed, the regions that would function best as species specific probes under conditions of

hybridization were not obvious. Extensive analysis of each 28S sequence yielded several potential probe sites. These were studied in detail to enable the selection of optimal unique sites for each probe, based on the need to obtain optimal hybridization characteristics under the test conditions. The highly specific hybridization characteristics of all probe sequences developed by us were then validated by experimental results. The prior existence in GenBank of sequences for Candida albicans and serotypes A and B (GenBank accession numbers L14067 and L14068) of Cryptococcus neoformans 28S genes was in itself not sufficient to enable even an individual versed in this field to develop specific probes for either of these two organisms. We had to obtain novel 28S sequence from Candida albicans, Candida glabrata, Candida guilliermondii, Candida kefyr, Candida krusei, Candida lusitaniae, Candida parapsilosis, Candida tropicalis, Cryptococcus neoformans serotype A. Cryptococcus neoformans var. gattii serotype B, Cryptococcus terreus, Cryptococcus laurentii, Filobasidiella (Cryptococcus) neoformans var bacillispora serotype C, Filobasidiella (Cryptococcus) neoformans var neoformans serotype D, Filobasidium 15 capsuligenum and Filobasidium uniguttulatum before we were able to identify potential regions for the development of species specific probes for these two fungal organisms that would not cross react with the others listed above.

Our modification of the Chomczynski technique (see Example 2, below) allows us to obtain DNA from any clinical specimen, irrespective of source (see Table 10 for a variety of clinical specimens tested), within a 3 hour period. The PCR amplification and subsequent probing can be accomplished with ease within a 24 hour period. The final identification is therefore possible in a day as opposed to several days or weeks required by traditional methods. This speed and sensitivity of diagnosis can make a difference between life and death in debilitated patients battling fungal diseases of undetermined cause. Rapid diagnosis will allow physicians to immediately direct their therapy towards curing the identified causative fungus, rather than wait for days or weeks while the patient succumbs to an unknown fungus.

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Our probes have the ability to pick out the correct target organism even in a mixed fungal infection because of their high level of specificity. The methods of Hopfer et. al. and Maiwald et. al., do not allow identification of individual species in a mixed fungal infection because restriction fragment length polymorphism results are

nearly impossible to interpret when multiple organisms contribute to the restriction fragments. Their method can therefore only be used on a pure culture, and this also does not save any diagnostic time, because the fungus first has to be grown in culture.

The probes developed by us allow rapid species identification of a large number of pathogenic fungi by using multiple probes against only one PCR amplified fragment of DNA. Coupled with our modified DNA extraction technique and our ability to accurately diagnose in the case of mixed organisms, this strategy can provide the greatest amount of diagnostic information in the shortest amount of time. This diagnostic strategy is also amenable to automation, which can result in even greater savings in time, money and effort.

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The sequences and the complement of the sequences claimed in this disclosure, along with any modifications to these sequences, may potentially be utilized in assays for the identification of fungi based on several existing methodologies, as well as future improvements and alterations of this technology. These techniques include, but are not limited to, assays based on hybridization, ligation, polymerization, depolymerization, sequencing, chemical degradation, enzymatic digestion, electrophoresis, chromatography and amplification. Furthermore, all such variations ultimately are based in some selection or amplification process, some ligand or some nucleic acid moiety that recognizes or utilizes the sequences (SEQ ID NO: 1) to (SEQ ID NO:23), (SEQ ID NO:75) and (SEQ ID NO:76) claimed in this application. Such variations include but are not limited to use of a variety of linear or exponential target amplification schemes, such as, any of the myriad forms of PCR, the ligase chain reaction, Q-beta repliase, etc.; direct detection of species-specific nucleic acid purified or extracted from pure fungal culture using a probe selected from the group (SEQ ID NO: 3) to (SEQ ID NO: 23), (SEQ ID NO:75) and (SEQ ID NO:76); use of the complementary DNA forms of (SEQ ID NO:1) to (SEQ ID NO:23), (SEQ ID NO:75) and (SEQ ID NO:76); use of the RNA forms of these sequences and their complements; and use of derivatives of these DNA or RNA sequences by the addition of one or more reporter moieties from a variety of labels including nucleic acid sequences, proteins, signal generating ligands such as acridinium esters, and/or paramagnetic particles. These techniques may be

utilized with DNA, RNA or modified derivatives used as either the target or the detection molecule.

In addition to the 25 sequences SEQ ID NO: 1 to SEQ ID NO: 23, SEQ ID NO:75 and SEQ ID NO:76, we also describe an additional 53 sequences SEQ ID NO: 24 to SEQ ID NO: 74, SEQ ID NO:77 and SEQ ID NO:78. These 53 sequences are inclusive of SEQ ID NO: 3 to SEQ ID NO: 23, SEQ ID NO:75 and SEQ ID NO:76 and are shown as a multiple sequence alignment (Table 3) with coordinate 1 corresponding to base # 431 of a reference *S. cerevisiae* 28S rRNA gene. (The numbers are comparable to the primary sequence of *S. cerevisiae* 28S rRNA gene. Genbank accession number: J01355). These sequences were obtained by amplifying and sequencing 28S rDNA from various fungi with primers SEQ ID NO: 1 and SEQ ID NO: 2. (SEQ ID NO: 1 corresponds to coordinates 403-422 and the SEQ ID NO: 2 corresponds to coordinates 645-662 of the reference *S. cerevisiae* gene).

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An analysis of these aligned sequences enabled us to develop the species specific probes SEQ ID NO: 3 to SEQ ID NO: 23, SEQ ID NO:75 and SEQ ID NO:76 and sites for these probes are shown underlined. These 53 aligned sequences contain sufficient variability, to enable a person versed in this art, to develop additional species specific hybridization probes in the 10-50 nucleotide length. Similarly, longer species specific hybridization probes encompassing the entire 200+ nucleotide length can also be envisioned. Species identification may also be accomplished by direct DNA sequence determination of any DNA amplified with primers SEQ ID NO: 1 and SEQ ID NO: 2. If the derived sequence matches approximately 98% or more of any sequence in SEQ ID NO: 24 to SEQ ID NO: 74, SEQ ID NO:77 and SEQ ID NO:78. then the identity of the organism can be ascertained. Additionally, we recognize that parts of SEQ ID NO: 24 to SEQ ID NO: 74, SEQ ID NO:77 and SEQ ID NO:78 may be specific for groups of fungi arranged phylogenetically at the level of genus or higher. SEQ ID NO: 24 to SEQ ID NO: 74, SEQ ID NO:77 and SEQ ID NO:78, their complements, along with any modification to these sequences may also potentially be utilized in assays for the identification of fungi based on existing methodologies and future technologies as noted above for SEQ ID NO: 1 to SEQ ID NO: 23, SEQ ID NO:75 and SEQ ID NO:76.

Table 3:

Multiple sequence alignment for (SEQ ID NO: 24) through (SEQ ID NO: 74), (SEQ ID NO: 77) and (SEQ ID NO: 78)

```
(RILIZO2) AGCCAGACTG GCTTGTCTGT AATCANTCTA GGCTTCG.GC CTGGATGCAC TTGCAGGCTA ..TGCCTGCC
(Rhizo3) AGCCAGACIG GCTTGTCTGT AATCAGTCTA AGCTTGG.GC TTGGATGCAC TTGCAGGCTA ..TGCCTGCC
[Rhizol] AGCCAGACIG GCITGICIGI AATCAATCIA GGTTTCGTGC CTGGATGCAC TTGCAGACIA TTTGCCTGC
(Mucor_) AGCCAGACTG GTTTGACTGT AATCAACCTA GAATTCGTTC .TGGGTGCAC TTGCAGTCTA ...TACCTGCC
(C Terr) AGTCAGTCAT GTCTATTGGA CTCAGCOGGT TCT.....G COGGTGTACT TCCTTTAGAT GGGGTCAAC.
(F_Caps) AGTCAGTCAT GTCTATTGGA CTCAGCOGT TCT.....G COGGTGTATT TCCTTTAGAT GGGGTCAAC.
(F Unig) AGTCAGTOGT GCTCAATOGA CTCAGOOG. TTC....T GOGGTGTATT TOCATTGGGT GGGGTCAAC.
(C Neob) AGTCAGTOST GTCTATTGGG TTCAGOCAGC TCT.....G CTGGTGTATT COCTITAGA. CGGGTCAAC.
(F Neoc) AGTCAGTOGT GTCTATTGGG TTCAGOCAGC TCT.....G CTGGTGTATT COCTTTAGA. CGGGTCAAC.
(C Neof) AGTCAGTOGT GTCTATTGGG TTCAGCCAGT TCT.....G CTGGTGTATT CCCTTTAGA. CGGGTCAAC.
(T Beig) AGTCAGTOGT GTTCTTTGGA TTCAGOCAGT TCT.....G CTGGTCTACT TOCTTGGAA. CGGGTCAAC.
(C Laur) AGTCAGTOGT GTCTGGGAGG CTCAGCOGGT TCT......G CCGGTGTATT CCTCTCAGA. CGGGTCAAC.
(Beauve) GACCAGACTT GGGCTTGGTT GATCATCOGG GGTTC.TCC. COGGTGCACT CTTCC.GGCC CAGGCCTACC.
[Fusari] GACCAGACIT GGGCTTGGIT AATCATCIGG GGITC.TCY. CCAGTGCACT TITCC.AGTC CAGGCCCACC.
(Acremo) GACCAGACTT GGGCTCGGTG AATCATCCGG CGTTC.TCG. CCGGTGCACT TTGCC.GTCC CAGGCCACC.
(Paccil) GACCAGACTI GGGCCCGGTG GATCATOCAG CGTTC.TCG. CTGGTGCACT CCGCCGGTT CAGGCCAGC.
(P_Boyd) GACCACACTT GIGCOCGTOG AATCAGCOCC COCTOGTOG. GCGCCCACT TOGGCCGCCT CAGGCCACC.
(S Brum) GROCAGACTO GOGOCOGTOG GATCAGOOGT OCCTOGTOG. GOGGOCOGACT OCCGOCOGCT OCCGOCOGACC.
(S_Brev) GACCAGACIT GOCCOCTOG GATCAACOCT COCTIG.CG. GOCCOCACT COCCOCGCT CACCOCACC.
(Sporot) GACCAGACIT GOSCOYCOGG GACCACOGGG CAGGCCACO CAGGCCACC.
(B Delm) GACAGAGIC GEOGRAGES GITCAGOGG CATTOST. TG COCCIGCACT COORTAGES CEGGOCACE.
(H_Caps) GAYCAGAGIC GEOGRAGGE GITCAGOGGE CATTOST.TG COCCEGNAT COCCEGNAT COCCEGNAT COCCEGNAT.
(A Nidu) GACCAGACTO GEOCOC.GGG GITCAROCAG CACTOG..TG CTGGTGTACT TOCOCGGGG CGGGCCAGC.
(A Ungu) GACCAGACTC GEOCTC.GGG GTTCAGCCAG CACTCG..TG CTGGTGTACT TCCCCGGGG CGGGCCAGC.
(A Ustu) GACCAGACTO GROCCIOG GITCAGOCAG CACTOG...TG CIGGIGITACT TOCOGGGGG CGGGCCAGC.
[A_CLAV] GROUNGICE GOTOGO.GGG GITCHGOOGG CATTOG...TG COGGIGERACT TOCOGGIGEG CGGGOODAGC.
(A Fimi) GRONGROTO COMONIGOS CTICAGOSS CATTOS...TG COSSIGNACT TOMOSTOSS COSSONASC.
(A Flav) GROCHGROTO COSTOC.AGG GITCHGOOGG CATTOG..TG COGGIGIACT TOCCTGGGGG CGGGCCAGC.
[A Ochr) CACCACACTE COCCC.CCC CITCACCCC CATTOC..TC COCCTGTACT TOCCCCCCC.
(A Nige) CACACACTO COMOC.COG CITCACOGG CATTOG..TG COGGIGIACT TOMOGIGGG COGGONAGE.
 (A Terr) AACCAGACTC GCTGGC.GGG GTTCAGGGGG GCTTGG..GC GGGGTGTACT TGGGGGGG GGGGGCAGC.
 (A_GLau) GACCAGACTO COTTOC.GGG GTTCAGCOGG CITTOG..GG COGGIGIACT TOCCGGGGGG CGGGCCAGC.
 (Penici) GACCACACTO COCCALGOS GITCACCOS CATTOS... TO COSSIGNACT TOCCASCOS COSSICACE.
 (C Immi) AMOCAGACTO GGTOGTOGGG GCTOGAGGGG CATGAGT.GC COGTGTACTC COCCAGGAGG.
 (Bipola) ACCAGACIT COTTCAGIT COTCATOCG COTTT.T.CC COGGGCACI CITATICAGG CAGGCCACC.
 (CHIVIL) ACCRETATE CONTRACT CONTRACT CONTRACT CHARGES CACCORACE.
 (Chryso) Alocadati Gososocc Gaicaicoc Teitc. T.CA cossiscaet coscosisci Cassocaec.
 (Clades) Anoxicacit ecrossest. Gitcoscos Tatia.t. CA cossicitaci ascoscitis chissocras.
 [Kalbua] Rescriptor execocoses ecitationes latifatale commence commence commence.
 [C Paix] CATCHCACT CONTINUE AIG..THAT CHOICEGG. .. GIGGOCTC TACHGITTAC COGGOCAGC.
 (C Imp) Gatchesiche Genäthiger aug.: Itach Tentosses. ... Giesoche Pacagintat Geografie.
 (CAIDI) GATCAGACIT GGIATITICC ATG. CICCI CICIOGGG. ..GOGGOGG TGOGGITTAC CGGGOCAGC.
 (CO11) CATCACACTO CATATITATO CACCOTTOC TROCTOCOS. .. GOGREGO COCACCITAT COCGOCIACO.
 (C Clab) GATCHCAT GENERATICS GOOGLIGON TOTOGROSS TRESCHOTOT CHARGICAL TEGGORAGE.
 (5 Cere) GATCAGACAT GETCHTTGT GOOTICIGCT OCHTGIGGGT AGGGGAATCT CGCATTTCAC TGGGOCAGC.
 (C Kefy) CATCHGAT GOOTITICAT
 (Geothi) NATONGACIT GETECTET...TETTCARCT RIGITIOGSC ATAGTETACT CAGCAGIACT AGGCCAAGG.
 {C Insi} AAGCAGACAC GGT......TITIAC CGGGCCAGC.
 (C Krus) COCCERCAT COSCATIONS CACCECTON TOTOGROSSE ..GOOGSTOT COGGITTOCC TECGOCACE.
 (Blasch) .....
 (P_Braz) ACCAGAGTCG GCCGCGGGG CTCAGCGGCC ACTCGT.TGC CCGTGCACTCCCCGTG....GTCGGGCCA
 (Pc_Hum) ATCAGACATG CCTTTATAGG AGATGCCATT GTT..TCGGC ATTGGCAGTA TTATCCGAAT TGGCAGGCCA
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	71
(Shizo2)	ANCONDIATT TOACTICAGE CANNAMACTA COCCONATOT COCC CACTTOTOGG TOTTATAGTC
(Rluzo3)	ANCCACANT TEGETTENES GANAMACTA AGGGANATET CCCC CATCOGTOGG TETTATAGTC
(Rhizol)	NACCACAATT TITTITGAGT GIANNAACIA TIGGANATGT GGCCAATATT TATTIATIGG TGTTATAGTC
(Mucor)	AACAACAGTT TGATTTGGAG GAAAAAATTA GTAGGAATGT AGCCTCTCGA GGTGTTATAG
(C Terr)	.ATCAGTTTT .GATCCCTGG AAAAGGGCAG GAGGAATGTA GCACTC.TCG GGTGAACTTA TAGCCTTCTG
[F Caps]	.ATCAGTITI .GACCGTTGG ATAAAGGCAG GAAGAATGTA GCACTC.TCG GGTGAACTTA TAGCTTCTTG
(F Unig)	.ATCAGTITI .GATCCCTGG ATAAAGGCAG GAGGAATGTA GCACCC.CCG GGTGAACTTA TAGCCTCTTG
(C Neob)	ATCACTICT -CATCCCTCC ATANCCCCTC CACCANTCTC CCACTCTTCC CCCTCTTA TACCCTCCTC
(F Neoc)	ATCAGITCI . CATCCGICG ATAACCCCIC CACCAATCIC CCACTCTICG CCGIGIGITA TACCCICCIC
(F Neod)	ATCAGITCT GATCGGTGG ATANGGGCTG GAGGAATGTG GCACTCTTCG GGGTGTGTTA TAGCCTCCTG
(C Neof)	ATCACTOC ATTOCOM ATAMOSCO COCCANTICIA CONCICTOS CACTOTOM TACOCTOMO
(T Beig)	ATCAGTITI GROOGTOG ATAMAGGIAG TAGGAATGTG ACTICTOC GGAAGTGTTA TAGCCTATTA
(C Laur)	.ATCAGTITT .GTCCCACGG ATAATCCCCG CGCCAAAGTA CCACCTCC GGGTGTGTTA TAGCCCCCTG
(Beauve)	ATCAGTICG CCCT.GGGGG ACAAAGCCTT CGGGAACGTG CCTCTCTCC
(Fusari)	ATCAGITIT COCC.GGGG ATAMARCIT CGGGAATGIG GCICYCYC
(Acremo)	ATCAGTTOG CCCC.GGGG ATAMAGGTTT CGGGAATGTA CCTCCTTC
(Paecil)	ATCAGTTOG COGC.GGGG AMAMAGGCTT CGGGAACGTG GCTCCTAC
(P Boyd)	ATCAGTICG CTGCAGGGG ACAAAGGCGA TGGGAATGTG GCTC. TTCGGA
(S Brum)	ATCAGITOG OCTOGGGGG AGAAAGGOGG OGGGAATGTG GCTC. TAC
(S.Brev)	ATCAGITOG .TOCCGCGCG AGAAAGCCCG CGGGAATGIG GCTCTTCGGA
(Sporot)	ATOGETICT C. OCAGGGG ACAMAGGCCG CGGGAACGTA GCTCCTTCG
(B Derm)	GROSSITIC GROSSOSS TOWNSCOOK COGGNATISTS TOSCOTOTO
(E Caps)	GROSSITIC GROSSOCS TOWNSCOOL COSCRATISTS TOSCCIOICGSGG.C
(A Nidu)	GROGGITTG GEOCOCCE TCANAGECCC CAGGRATGIA TOCCCTCCGEGGIT
(A Ungu)	GTOSTITE GGOGGOOG TCAAAGCOC CAGGAATGTA TCACCTCCGGGGTT
(A Ustu)	CTOSCTTT: GCCCCCCCC TCAAAGGCCC CAGGAATGTG TCGCCCTCCGGGG.C
(A Clav)	GTOCCTITG GGCCCCCCC TCAAACCCCT CCCCAATGTA TCACCTCTCGGGG.T
(A Fumi)	GTOGGTTTG .GGGGGOOGG TCHANGGOOG TOGGANGTA TCHOCTCTC GGGG-T
(A Flav)	GROSCITTG GGOGGOGG TCHANGGCTC COGGNATGIA GIGCOCTYCGGGG.C
(A Ochr)	GROGGITTG GGCGGCCG TCAAAGGCCC CCGGAIGIA GCACCCITCGGGG.I
(A Nige)	GROGGITTG .GGCGCCCG TCAAAGCCCC CTGGAATGTA GTROCCTCCGGGG.Y
(A Terr)	GTOGGTTTG GGOGGOOGG TCAAAGGOCT COGGAATGTA GOGOOCITCGGGG.C
(A Glau)	GTOSCTTTG GGCCCCCG TCAAAGGCCC CTGGAATGTA ACGCCTCTCGGGG.C
(Penici)	CHARLETTE GROCOCC TOAAACCOC TOCAATRIA ACCOCCCCGGG.C
(C Immi)	ATCACTICT GOOGGTTGG TTAAAGGOOT CTGGAATGTA TOGTOCTOCGGGAC
(Bipola)	ATTACTUTE COOCCIGG ATARAGGICT CIGTCACGIA CCITOCITCGGGIIG
{Curvul}	ATTACTURE GEOGGICGG ATAMAGGICT CIGACACGIT CCITCCITCGGGITG
(Chryso)	ATTOCHTET CONCENTRE ATAMAGEOOC TAGGAATGIG GCTOCTCICGGGGAG
(Clades)	ATTOMORE TO TOTAL AT ARCHOTT CACCALIGIA COTOCTOGCCACIG
	GGG-BC
CO Damet	REVINCTION CROCCETEC CETTARCTICA ARGARATGEG GCACIGATIC
10 mm1	- Band Color Cococcus Cacally Car Tilly Alleig Grandollo Galley
(C Albi)	RECOGNEY CHOCOCKE CHARACTEC CACCATTEG CCACCATUC LOUIGE
(c ent)	ATTOCKET COCCUCTAG CATAATGGG TAGGAATGTG ACTITICATICGGIGGA
(0.01-1-1	ANOSCITT G.GOGGOG GAMAMACT AGGGANGIG GCICIGGGC TOGGIGIAGA
(S Carrol	ATVACTITTI G. GTEGCAG GATANATOCA TAGGANTGTA GCITGCCICGGTAN
(C Kaferi	ATCACTITT AGGGGGG GATANATOCA TAGGAATGTA GCTTGCCTC GGTAA ATCACTITT AGGGGTG GATANATOCT CGGGAATGTG GCTCGCCTTC GGTAGA
(Cartery)	TODOCTOCTOR TOCOGRAP CANANGANG TAGGRAGGIA ROLLILO
(C Insi)	CONTRACT CONTRACT CAGARTGIG GOUGHOUT 100010000
_	A AMERICAN ACCUPATION ACCUPATION OF THE CONTROL OF
(C_Krus)	
(Blasch	TOTAL CONTRACTOR CONTR
(P_Braz) GCGTCGGTTT CGACGGCCGG TCAAAGGCCC CCGGAATGTGTCGCCTCTCGGGG
" (Pc_Hum) GCATCGGTTT CACTTACTGG ATAAAACTGG AAGAAGGTAGGCTCTCTTCGGAGGGTTTTT TAGCTTCCAG

	141	210
(Nuizo2)	141 CCTTAGAAAA TACCTIGGGT TGGATTGAGG AACGCAGCGA ATG	CITATTG
(Nuizo2)	CCTTAGABAA TACCTTGGGC TGGATTGAGG TACGCAGGA ATG	CIATTIG
(Rhizol)	CTTTAGABAA TACCTTGAAT TGGATTGAGG AACGCAGCGA ATGCTTCTCT TTAGAGGCAA	AGICITTIAT
{Mucor }	CCTACTATCA TACTCTGGAT TGGACTGAGG AACGCAGCGA ATGCCATTAG GCRAGATTGC	TGGGTGCTTT
{C Terr}	TYCTETECE TYCTTGGGAC TGAGGAAGGC AGCATGCCTT TATGGCCGGG GTTCGCCCAC	GTACATGCTT
(F_Caps)	TOPOTACE TOCHTOCOGO TORGONOGO AGCATGOCTT TATGGOOGGG ATTOCTOCAC	GTACATGCTT
(F Unig)	TCACATACAG TGGTTGGGAC TGAGGAACGC AGCATGCCTT TATGGCCGGG ATTCGTCCAC	GIACATGCIT
(C Neob)	TOGOTHOR TOGTTGGGAC TGAGGAATGC AGCTGGCTT TATGGGGGG GTTGGGCAC	GITOCAGCIT
(F Neoc)	TOTATACAC TOTATOGGAC TOTAGGAATGC AGCTOGCCIT TATGGCOGGG GITOGCCAC	GITOGAGCIT
(F Neod)	TOTAL TOTAL TOTAL TOTAL TOTAL TOTAL ACCIDENCE TATESCOOR GITCHOOCK	GITOGAGCIT
(C Neof)	TOGETHERE TECTTOGGAC TGAGGATGC AGCTOGOCIT TATGGOOGG GITOGOCCAC	GITGGAGCIT
(T Beig)	THE THE TEXT TEXT AGAC TEAGGACTEC ACTIOSOCIT TAIGCCOCC CTTOSGCAC	GITOGAGCIT
(C Laur)	TOTATION CONTRACAC TORGOCATOR ACCTOSOCIT TATGCCAGG GITOCOCCAC	TITOGAGCIT
(Beauve)	G TOTTATAGOC COTTGOGTAA TACC_CIGIG GOSGACIGAG GITCGOG	CATTOGCA
(Fusari)	G TOTTATAGOC COTTGYGTAA TACC.CTGGB GGGGACTGAG GTTCGCG	CWICIGCA
(Acremo)	C TOTTATACOC OCTIGOCIAA TAOC.CIGGC GIGGACIGAG GIOCOCC	C.TCIGCA
{Paecil}	G TOTTATAGOO OGTIGOATAA TAOO.CIGGG GOGGACIGAG GITOGOG	C.TOUSCA
(P Boyd)	C TYTTATAGOC OGOGGGGAA TAOC.OCTOG GOGGACTGAG GACCECE	CATCIGCA
(S Brum)	G TYTTATAGOC OGOOGOGIAA TAOC.COOG GOGGACTGAG GACOGOG	- CONTACOLA
(S Brev)	C TETTITAGOC COCCEGIA TACO.CIOSE GIGGACIGAE GACOSOG	CGIAIGCA
(Sporot)	C TOTALIACOC COCCOCCA TOCCLOCICE GEGENOCIAE GACCECE	CITOGGCA
(B Derm)	C TOTALIA COL GEGGIGCAA TOCGOCAGI CGGACCEAG GAACCE	CITUGGA
(H Caps)	C TOTTATAGOO GEGGGIGCHA TEOGGOOGHGI OSGGAOOGAG GAACGOG	CICUGUA
(A Nidu)	C TETTATACOC TOCCOTOCAL TOCCOCCACO COGRACORA CALCOCA	Critusua
(A Ungu)	C TOTATAGOC TOGGETGCAA TGOGGOCAGC CTGGAOOGAG GAACGCG	CIIUGGA
(A Ustu)	G TOTTATAGOC TEGGGTGCAA TGOGGOCAGC COGGACOGAG GAACGOG.	CIICGCA
[A Clav]	G TOTTATAGOO GOGGETGCAA TGOGGOOTIGO CTGGACOGAG GAACGOG	
[A Fumi]	G TOTTATAGOO CAGGGTGCAA TGOGGOOTIGO CTGGAOOGAG CAACCOG.	
(A Flav)	A COTTATAGOC GOGAGIGCAA TGOGGOCAGC CTGGACOGAG GAACGOG.	
(A Ochr)	G COTTATAGOC GEGGGTECHA TECCGOCAGC CTGGACCGAG CAACCCG.	
(A Nige)	A COTTATAGOC AGGGGTGCAA TGCGGCCAGC CTGGACCGAG CAACGCG.	
(A Terr)	G COTTATAGOC GEGEGTECHA TECCECCACC CTGGACCEAE CAACCCG.	CTTOGGCA
(A Glau)	G CTIATAGOC AGGGTGTVA TGOGGCAGC CTGGACCGAG GAACGCG.	CTTOGGCT
(Penici)	G TOTTATAGOC GAGGIGOCA TGOGGOCAGC HCAGACGGAG GAACGCG.	CTTOGGCA
{C_Inmi}	G TOTTATAGOC AGGGGGGGAA TGCGGCCAGC CGGGACTGAG GAACGCG.	CATCTCCT
(Bipola)	G CONTATAG. G GENERACTION TROCHOCAGO CTOSECTICAG GTOCOCG.	CATCTCCT
(Corvor)	G CATATAG. G GEAGAGGICA TACCACCAGC CIGGACTGAG GICCOGG.	CITIOGGCI
(Chryso)	TANTA C CONCILICAN ICCHECANCE COCCESCAR CHOCCE. G CANADAR C CONCILICAN ICCHECANCE CHOCCE.	CITOGGCI
(Clados)	TIATA G CHICHIGIGA TOCARCARO GORROLLO GIORDO	CTTOGGCA
{Malbra	G TOTTATAGO AAGGGTGCAA TGCGGCCAGC CGGGGCTGAG GAACGCG.	G CITOG.GOOT
(C_Para)	G TCTTATAGO: MAGGAGAA TGCCCAGC TTACACTGAG GACTGOG. G TGTTATAGO: T.TTGTC.GA TACTGOCAGC CTACACTGAG GACTGOG.	G TITAT ACCT
(C_Irop	G TGTTATAGOC T. CTCAC. CA TECTGOCAGC CTACACOCAG CACTGOG.	G TITITAACCT
(C'YPPI)	G TGTTATAGOC T-CTGGC-GA TGY-GOCAG CACTGGG-	A TITT ATCA
(c ⁻ enn	G TETTATAGO: T.COGIT.GA TECTEOCIGE CIRCACOGAG GACTECOG. G TETTATAGOE C. TEGGG. AA TACGOOCAG CECTECOGA G TETTATAGOE C. TEGGG. AA TACGOOCAG CACTECOGA COMPANYON TO THE COMPANYON THE COMPANYON TO THE COMPANYON TO THE COMPANYON TO THE COMPANYON	TA CITCITATOI
(C_@mp	G TGTTATAGOC C. TGGGG.AA TALOGGCAGC TGCTGCAGC GACTGGGA	OG TAAGTCA
(S_Cere	G TATTATAGOC T. GTGGG. AA TACTGOCAGC TGGGACTGAG GACTGOGA	CT TITGICA
	G TATTATAGOC T. GIGGGAAA TACAGOCAGO TOGGACTGAG GATTGOGA G TGTTATAGOC T. ACTIT. CA TAGCTOCTCA GGOGOCTCAG GACTGOG. C TGTTATAGOC T. ACTIT. CA TAGCTOCTCA GGOGOCTCAG GACTGOG.	
(Geotri	G TGTTATAGOC T. ACTITICA TARACTURA CONTINUAS GOCIGOS	T. TCT
(C_Krus	G TGTTATAGOC A.GGGOCAT TGCTGGGATA AGAGGCTGGG GTTTGAAA	TA ATTGTTTTC
(Blasch	TGAAA TIGITGAAAG GOAAGOCOAL GGLAGGALAA TAGAGGALA	ייי רנפפרא
(P_Braz	2)CG TCTTATAGCC GGGGGTGCAA TGCGGCCAGT CGGGACCGAG GAACGCG	
(Pc_Huπ	TAGCTGCAGT GACCGGGACC GGAAGGGAAA TTGGGTCTTT GAAGACCTTA TGATGTTC	GC AGAAATGGTC

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250
        211
        GCGAGTTTTC CAGGAAGGT. ....TTTCT GAGGTACTAC
                                                   SEQ ID NO: 68
(Rhizo2)
(Rhizo3) GCGAGTTGGC TGGGAATAT. ....TTTCT GAGGTGCTTT
                                                   SEQ ID NO: 69
(Rhizol) TGGGATTTAC GGATCAGAC. ....TGTGG CATTGTCACA
                                                   SEQ ID NO: 67
(Mucor_) CGCTAATAAA TGTTAGAATT TCTGCTTCGG GTGGTGCTAA
                                                   SEQ ID NO: 63
[C_Terr] AGG..ATGTT GACATAATGG CTTTAAACGA CCCGTCTTGA
                                                   SEQ ID NO: 53
        AGG..ATGIT GACATAATGG CITTAAAGGA COOGTCITGA
                                                   SEQ ID NO: 56
(F Caps)
        AGG..ATGIT GACATAATGG CTTTAAACGA CCCGTCTTGA
                                                   SEQ ID NO: 59
(F Unig)
        AGG..ATGTT GACAAAATGG CTTTAAACGA COOGTCTTGA
                                                   SEQ ID NO: 50
(C Neob)
        AGG. ATGIT GACAAAATGG CITTAAAOGA COOGICITGA
                                                   SEQ ID NO: 57
(F Neoc)
        AGG. .ATGIT GACAAAATGG CITTAAACGA COCGTCITGA
                                                   SEQ ID NO: 58
[F Neod]
        AGG..ATGIT GACAAAATGG CITTAAACGA COCGTCITGA
                                                   SEQ ID NO: 51
(C Neof)
        AGG..ATGIT GACATAAIGG CITTAAACGA COOGICIIGA
                                                   SEQ ID NO: 74
(T Beig)
        AGG..ATGIT GAOGIAATGG CITTAAACGA COOGICITGA
                                                   SEQ ID NO: 48
{C.Laur}
(Beauve)
        AGG..ATGCT GGOGTAATGG TCATCAGTGA COOGTCT...
                                                   SEQ ID NO: 35
(Fusari) AGG..ATGCT GGCCTAATGG TCATCAACGA CCCGTCTTGA
                                                   SEQ ID NO: 55
(Acremo) AGG..ATGCT GGOGTAATGG TCATCAGTGA COOGTCTTGA
                                                   SEQ ID NO: 24
SEQ ID NO: 64
        AGG..ATGCT GGGGTAATGG TCATCAGGGA COOGTCTTGA
{Paecil}
(P Boyd) AGG. ATGCT GGCGTAATGG TOGTCAGOGA COCGTCTTGA
                                                    SEQ ID NO: 66
                                                    SEQ ID NO: 72
        AGG. ATGCT GGGTRATGG TOSTCAGOGA COOSTCTTGA
(S Brum)
                                                   SEQ ID NO: 71
        AGG..ATGCT GGOGTAATGG TOGTCAGOGA COOGTCITGA
(S Brev)
        AGG..ATGCT GGOGTAATGG TCACCAGOGA ACOGTCTTGA
                                                   SEQ ID NO: 70
(Sporot)
        CGG.. ACCCT GGCTTAATGG TOGTAAGOGA COOGTCTTGA
                                                   SEQ ID NO: 38
(B Derm)
        CGG...ACCCT GCCTTAATGG TOGTCAGOGA COOGTCTTGA
                                                   SEQ ID NO: 61
(H Caps)
(A Nidu) CGG. ACCCT GGOGTAATGG TOGCAAACGA COCGTCTTGA
                                                   SEQ ID NO: 29
(A Ungu) OGG...ACCCT GGCATAATGG TTGCAAACGA CCCGTCTTGA
                                                   SEO ID NO: 33
(A_Ustu) CGG...ACCCT GGGGTAATGG TOGCAAACGA COCGTCTTGA
                                                   SEQ ID NO: 34
(A CLAV) CCG...ACCCT GCCCTAATCG TOCTAAATCA CCCCTCTTCA
                                                   SEQ ID NO: 25
(A Fumi) CGG.. ACCCT GGCGTAATGG TOGTAAATGA CCCGTCTTGA
                                                   SEQ ID NO: 27
(A Flav) CCG.. ACCCT GCCATAATGG TOGYAAACGA COCGTCTTGA
                                                   SEQ ID NO: 26
(A Ochr) CGG..ACCCT GGCATAATGG TCGTAAACGA CCCGTCTTGA
                                                   SEO ID NO: 31
(A Nige) CGG.. ACCCT GGCATAATGG TCGTAAACGA CCCGTCTTGA
                                                   SEQ ID NO: 30
(A Terr) CGG. ACCCT GGCATAATGG TTGTAAACGA CCCGTCTTGA
                                                   SEQ ID NO: 32
         CGG. ACCCT GGCATAATGG TOGTAAAOGA COOGTCTTGA
                                                   SEQ ID NO: 28
{A Glau}
         CGG. ACCCT GGCATAATGG TOGTAAA...
                                                   SEQ ID NO: 65
{Penici}
         CGG...ATGCT GGCATAATGG TTGTAAGCGG CCCGTCTTGA
                                                   SEO ID NO: 45
{C Immi}
         AGG..ATGCT GEOGRAATGG CTGTAAGGGG CCCGTCTTGA
                                                   SEQ ID NO: 36
{Bipola}
         AGG. ATGCT GEOGRAPHICG CTGTAAGCGG COCGTCTTGA
                                                   SEQ ID NO: 41
{Curvul}
(Chryso) AGG..ATGCT GGOGTAATGG TIGTAAGGGG COGTCITGA
                                                   SEQ ID NO: 39
(Clados) AGG. ATGCT GGGTAATGG TOGTAATGCG COOGTCTTGA
                                                   SEQ ID NO: 40
(Malbra) CGG. ATCCT GCCGTAATGG CTGTAAGCGG CCCGTCTTGA
                                                   SEQ ID NO: 62
(C Paix) AGG..ATGIT GGCATAATGA TCTTAAGTGG CCCGTCTTGA
                                                   SEQ ID NO: 52
(C_Trop) AGG..ATGIT GGCATAATGA TCITAAGTGG CCCGTCLIGA
                                                    SEO ID NO: 54
                                                    SEQ ID NO: 42
(C Albi) AGG. ATGIT GGCATAATGA TCTTAAGTCG CCCGTCTTGA
(C Guil) AGG. ATTECT GECATAATEA TOOCAAACOG COOGTETTEA SEQ ID NO: 44
                                                    SEQ ID NO: 43
(C Glab) AGG. ATGCT GGCATAATGG TTATATGCCG COGTCTTGA
(S_Cere) AGG..ATGCT GGCATAATGG TTATATGCCG CCCGTCTTGA
                                                    SEQ ID NO: 73
{C Kefy} AGG..AIGCT GGGGIAAIGG TIAAAIGGGG GGGGCTTGA
                                                    SEQ ID NO: 46
                                                    SEQ ID NO: 60
 (Geotri) AGG. ACCIT GGCATAATCA TICTATACCG COCGTCTTGA
 (C_lusi) Agg. Aggr geographeg thechagog constitues
                                                    SEQ ID NO: 49
 (C Krus) CGG. ATGCT GSCAGAACGG CCCAACACGG CCCGTCTTGA SEQ ID NO: 47
 (Blasch) GOODCROGET CTCCTCRGCC TGCTTTCGCR CCCGTCTTCR SEQ ID
                                                            NO:
                                                                  37
(P_Braz) CGG..ACGCT GGCTTAATGG TCGTAAGCGA CCCGTCTTG- SEQ ID NO: 77
 (Pc_Hum) CTAAGCGACC CGTCTTGAAA CACGGACCAA GGAGTCTAAT SEQ ID NO: 78
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Legend to Table 3:

The multiple sequence alignment shows the sequence of 28S ribosomal RNA genes amplified with primers SEQ ID NO: 1 and SEQ ID NO: 2. 23 species specific probes (SEQ ID NO: 3 to SEQ ID NO: 23, SEQ ID NO:75 and SEQ ID NO:76) are shown underlined. Minor sequence variation among two isolate of the same organism are represented by the appropriate code (see key below). Major differences among *Rhizopus* species are depicted by including 3 separate *Rhizopus* sequences in the alignment. (The organisms in this figure are listed according to their sequence relatedness.)

Key to symbols:

- (.) gap in sequence to facilitate alignment
- 15 (R) A or G

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- (W) A or T
- (Y) T or C
- (M) A or C
- (K) T or G
- 20 (S) G or C
 - (B) T,G or C

Acremo Acremonium species
A_clav Aspergillus clavatus

25 A_flav Aspergillus flavus

A_fumi Aspergillus fumigatus

A_glau Aspergillus glaucus

A_nidu Aspergillus nidulans

A_nige Aspergillus niger

30 A_ochr Aspergillus ochraceus

A_terr Aspergillus terreus

A_ungu Aspergillus unguis

	A_ustu	Aspergillus ustus
	Beauve	Beauveria species
	Bipola	Bipolaris species
	Blasch	Blastoschizomyces species
5	B_derm	Blastomyces dermatitidis
	Chryso	Chrysosporium species
	Clados	Cladosporium species
	Curvul	Curvularia species
	C_albi	Candida albicans
10	C_glab	Candida glabrata
	C_guil	Candida guilliermondii
	C_immi	Coccidioides immitis
	C_kefy	Candida kefyr
	C_krus	Candida krusei
15	C_laur	Cryptococcus laurentii
	C_lusi	Candida lusitaniae
	C_neob	Cryptococcus neoformans var gattii serotype B
	C_neof	Cryptococcus neoformans serotype A
	C_para	Candida parapsilosis
20	C_terr	Cryptococcus terreus
	C_trop	Candida tropicalis
	Fusari	Fusarium species
	F_caps	Filobasidium capsuligenum
	F_neoc	Filobasidiella (Cryptococcus) neoformans var bacillispora
25		serotype C
	F_neod	Filobasidiella (Cryptococcus) neoformans var neoformans serotype D
	F_unig	Filobasidium uniguttulatum
	Geotri	Geotrichum species
	H_caps	Histoplasma capsulatum
30	Malbra	Malbranchea species
	Mucor_	Mucor species
	Paecil	Paecilomyces species

Penici Penicillium species P boyd Pseudallescheria boydii P braz Paracoccidioides brasiliensis Pc-Hum Pneumocystis carinii, human isolate Rhizo1 Rhizopus species isolate #1 Rhizo2 Rhizopus species isolate #2 Rhizo3 Rhizopus species isolate #3 Sporot Sporothrix schenkii S brev Scopulariopsis brevicaulis S brum Scopulariopsis brumpti S cere Saccharomyces cerevisiae T beig Trichosporon beigelii Pneumocystis carinii

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Further variations of the invention that utilize any of the named sequences will be apparent to those with ordinary skill in the art. The following examples illustrate various aspects of the invention but are not intended to limit its usefulness.

EXAMPLE 1. Testing probes SEQ ID NO: 3 to SEQ ID NO: 23, SEQ ID NO:75 and SEQ ID NO:76 for hybridization specificity.

Probes listed in SEQ ID NO: 3 to SEQ ID NO: 23, SEQ ID NO:75 and SEQ ID NO:76 were tested for specificity against their target organisms. Probe SEQ ID NO: 5 for Candida albicans was the first one tested against a panel of fungi taken from the Mayo Clinic collection. 28S rDNA from Acremonium sp., Aspergillus clavatus, Aspergillus flavus, Aspergillus fumigatus, Aspergillus glaucus, Aspergillus nidulans, Aspergillus niger, Aspergillus ochraceus, Aspergillus terreus, Aspergillus unguis, Aspergillus ustus, Aspergillus sp., Beauvaria sp., Bipolaris sp., Blastomyces dermatitidis, Candida albicans, Candida glabrata, Candida guilliermondii, Candida kefyr, Candida krusei, Candida lusitaniae, Candida parapsilosis, Candida tropicalis, Chrysosporium sp., Cladosporium sp., Coccidioides immitis, Cryptococcus neoformans serotype A, Curvularia sp., Fusarium sp., Geotrichum sp., Histoplasma

capsulatum, Mucor sp., Penicillium sp., Pseudallescheria boydii, Rhizopus sp., Saccharomyces cerevisiae, Scopulariopsis brevicaulis, Sporothrix schenkii and Trichosporon beigelii was amplified in a polymerase chain reaction using oligonucleotide probes SEQ ID NO: 1 and SEQ ID NO: 2. All PCR amplifications were carried out as hot-start reactions in a 50 ul reaction volume using Perkin-Elmer (Norwalk, CT) 0.5 ml thin-wall polypropylene tubes and a Perkin-Elmer thermal cycler. Reagents added to the tube initially were 2.5 ul of 10X PCR buffer (100 mM tris pH 8.3, 500 mM KCl, 15 mM MgCl₂), 5.0 ul of 50% glycerol/1 mM cresol red, 8.0 ul of dNTP mix (1.25 mM each of dATP, dGTP, dTTP and dCTP), 11 picomoles of each nucleic acid primer and sterile water to make up a volume of 25 ul. A wax bead (Ampliwax Gem=100, Perkin-Elmer) was added and the tubes heated to 77°C for 30 seconds and cooled to room temperature to form a wax barrier. 2.5 ul of 10X PCR buffer, 5.0 ul of 50% glycerol/1 mM cresol red, 0.2 ul Taq polymerase (AmpliTaq 5U/ul, Perkin-Elmer) and 15.3 ul of sterile water was added to the tube along with 2.0 ul of DNA from the fungal whole cell boiled lysate described above. 50 cycles of thermal cycling was carried out at 94°C - 30 sec, 50°C - 1 min, 72°C - 2 min. Five microliters of polymerase chain reaction mix from each sample was run on a 5% polyacrylamide gel to visually confirm the successful amplification of 28S rDNA from each fungus listed above. 40 ul of the remaining amplified 28S rDNA was denatured in 1 N NaOH, and half of this denatured rDNA was slot blotted on to a positively charged polysulphone based membrane equilibrated in 0.5 N NaOH. The membrane was air dried for 15 minutes and baked in a vacuum oven at 80°C for 30 minutes. Amplified rDNA from each species was now bound and immobilized at a separate spot on the membrane. The free binding sites on the membrane were blocked by incubating the membrane for 3 hours at 40°C in hybridization buffer (100 ml of hybridization buffer was made using 1g non-fat milk powder, 6g NaH2PO4, 7g SDS, 200 ul 0.5M EDTA and adjusted to pH 7.2 with NaOH). The specific probe for Candida albicans (SEQ ID NO: 5) was end-labeled with radioactive phosphorus using 32P ATP and T4 polynucleotide kinase. 50 picomoles of this probe was added to 70 milliliters of hybridization buffer and the membrane was probed at 40°C overnight. The membrane was washed in hybridization buffer at 40°C for 15 minutes followed by a wash in 2X SSC at 40°C for 15 minutes. The membrane was then exposed on x-

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ray film for at least 1 hour. The oligonucleotide probe SEQ ID NO: 5 only hybridized to amplified 28S rDNA from *Candida albicans* (see Table 4) Under these hybridization conditions, probe SEQ ID NO: 5 is extremely specific for *Candida albicans*. The sequence of oligonucleotide probe SEQ ID NO: 5 differs from the sequences of other species of *Candida* by as few as 1 or 2 bases, but these mismatches are sufficient to prevent stable hybrids from forming with the other *Candida* species.

Probes SEQ ID NO: 3 to SEQ ID NO: 23 were tested for specificity, as described above for the *Candida albicans* probe SEQ ID NO: 5, against the same panel of fungi listed in the preceding paragraph. The positively charged polysulphone based membrane probed with *Candida albicans* probe SEQ ID NO: 5 was washed in 0.5 N NaOH at 40°C for 10 minutes to remove all bound *Candida albicans* probe. The membrane was sequentially probed with all probes listed in SEQ ID NO: 3 to SEQ ID NO: 23. For each subsequently tested probe, the membrane was blocked for at least 30 minutes, probe hybridization was carried out at 40-42°C for at least 3 hours, and post-hybridization washes were done in 2X SSC for 20 minutes. The membrane was stripped between probings by washing in 0.5 to 1.0 N NaOH at 40-42°C. Results are listed in Tables 4 to 7.

Probes SEQ ID NO:75 and SEQ ID NO:76 were tested for specificity against a modified panel of fungi, using the procedure just described. The specific organisms tested and the results obtained are listed in Table 8.

As shown in Tables 4 to 8, each probe listed in SEQ ID NO: 3 to SEQ ID NO: 23, SEQ ID NO:75 and SEQ ID NO:76 specifically hybridizes to only one target fungal 28S nucleic acid sequence. This specificity is essential for identifying a given species of fungus in clinical specimens containing mixed fungal organisms with a high level of reliability. The organisms listed in these Tables represent a majority of organisms that are commonly isolated from clinical samples. While we have developed 23 species specific probes (SEQ ID NO: 3 to SEQ ID NO: 23, SEQ ID NO:75 and SEQ ID NO: 76) that identify a total of 21 individual organisms, the additional organisms listed in the test panel were used to ensure that our probes did not have any cross-reactivity with other fungi likely to be present in a clinical specimen. The ability to accurately and reliably diagnose, and identify to a species level, this large a number of pathogens is unmatched by any other report. The fact that

we can achieve this by probing DNA amplified by a single pair of "Universal" probes (SEQ ID NO: 1 and SEQ ID NO: 2) is highly advantageous as it saves time, money and effort by providing the ability to test a single amplified target with 23 different probes (SEQ ID NO: 3 to SEQ ID NO: 23, SEQ ID NO:75 and SEQ ID NO:76).

A GenBank search was carried out with all probes listed in SEQ ID NO: 3 to SEQ ID NO: 23, SEQ ID NO:75 and SEQ ID NO:76 in order to determine whether similar gene sequences were present in the database. 28S sequences for Candida albicans Cryptococcus neoformans are already present in GenBank, and as expected, the probes for Candida albicans and Cryptococcus neoformans correctly identified the 28S sequences from these two organisms. Ten other probes also matched DNA sequences from a variety of genes not related to the 28S gene (Table 9). This was expected because short stretches of sequence identity can often be found for any query sequence in unrelated genes from the same or a different organism. This observation is known to those versed in this art. In all cases, sequences that matched a probe sequence were not located within the 28S rRNA genes. Our probes are used to analyze 28S DNA that has been previously amplified in a polymerase chain reaction with our probes SEQ ID NO: 1 and SEQ ID NO: 2. Under stringent conditions, these two probes only amplify DNA from fungal 28S rRNA genes. Therefore no amplified DNA from the non-28S genes listed in Table 9 will be available for the hybridization of probes SEQ ID NO: 3 to SEQ ID NO: 23. The presence of related sequences in non-28S, unamplified genes will not be detected and will, thus, not have any effect on the sensitivity or the specificity of our detection and identification strategy.

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Table 4:

Detection of species specific 28S sequence with probes SEQ ID NO: 3 to SEQ ID

NO: 8

FUNGUS	SEQ ID:	SEQ ID:	SEQ ID:	SEQ ID:	SEQ ID:	SEQ ID:
	3	4	5	6	7	8
Acremonium sp.	-	-	- .	-	-	-
Aspergillus clavatus	*	-	-	_	-	
Aspergillus flavus	-	•	-	-	. •	-
Aspergillus fumigatus	+	-	-	-	-	-
Aspergillus glaucus	-	•	-	-	-	-
Aspergillus nidulans	•	-	<u>-</u>	-	•	-
Aspergillus niger	-	-	-	-	-	-
Aspergillus ochraceus	-	. <u>-</u>	-	-	-	- ,
Aspergillus terreus	. -	-	-	-	-	-
Aspergillus unguis	-	-	-	-	-	. •
Aspergillus ustus	-	-	-	<u>-</u> ·	-	-
Aspergillus sp.	-	-	-	-	-	-
Beauvaria sp.	-	-	•	-	-	-
Bipolaris sp.	-	-	-	-	-	-
Blastomyces	-	+	-	-	-	•
dermatitidis				` .	:	
Candida albicans	-	-	+	-	-	-
Candida glabrata	-	-	•	-	-	-
Candida	-	-	-	-	-	-
guilliermondii			·			
Candida kefyr	-	-	-	-	-	-
Candida krusei	-		-	-	-	-
Candida lusitaniae	-	-	-	-	-	-
Candida parapsilosis	•	-	-	-	-	
Candida tropicalis	-	-	_	-	-	-

PCT/IB98/00865

		~,··				·
Chrysosporium sp.	•	-	• ,	-	•	-
Cladosporium sp.	-	-	-	-	-	-
Coccidioides immitis	-		-	+	-	-
Cryptococcus	•	-	-	 •	+	+
neoformans		į.				
Curvularia sp.	-	•	•	-	-	-
Fusarium sp.	-	-	-	-	-	-
Geotrichum sp.	-	-	-	-	-	-
Histoplasma	-	-	-	-	-	-
capsulatum						
Mucor sp.	-	-	-		-	-
Penicillium sp.	-	-	-	-	-	-
Pseudallescheria	-	-	-	-	-	-
boydii						
P. brasiliensis	· -	-		-	-	-
Pneumocystis carinii	-	-	-	-	-	-
Rhizopus sp.	-		•	-	-	-
Saccharomyces	-	-	-	-	-	<u>-</u>
cerevisiae						
Scopulariopsis		-	-	-	-	-
brevicaulis						
Sporothrix schenckii	-	-	-	-	-	-
Trichosporon beigelii	-	-	-	-	-	-
	1	1	3	l		l

+	Positive
-	Negative after 20 minute wash in 2X SSC

Table 5:

Detection of species specific 28S sequence with probes SEQ ID NO: 9 to SEQ ID

NO: 14

FUNGUS	SEQ ID:	SEQ ID:	SEQ ID:	SEQ ID:	SEQ ID:	SEQ ID:
	9	10	11	12	13	14
Acremonium sp.			_	-	-	
Aspergillus clavatus	-	-	_		_	· -
Aspergillus flavus	-	-	_	<u> </u>		
Aspergillus fumigatus	-	-	-		-	_
Aspergillus glaucus		+	_	•	•	
Aspergillus nidulans	-	-	-	-	-	_
Aspergillus niger	-	-	+		-	
Aspergillus ochraceus	-	•	-	-	_	-
Aspergillus terreus	-	-	· •	+	-	
Aspergillus unguis	•	-	-	-	-	- V
Aspergillus ustus	-	· -	-	-	-	-
Aspergillus sp.	•	•	-	-	-	-
Beauvaria sp.	-	-	-	-	-	*
Bipolaris sp.	-	-	-	-	-	-
Blastomyces	<u>-</u>	-	-	-	•	<u>-</u>
dermatitidis						
Candida albicans	-	· -	-	-	-	_
Candida glabrata	-	-	-	-	+	-
Candida	-	-	-	•		+
guilliermondii				;		
Candida kefyr	-	-	-	-	-	-
Candida krusei	-	-	-	-	-	-
Candida lusitaniae	-	-	-	-	-	•
Candida parapsilosis	<u>-</u>	-	-	-	. -	· -
Candida tropicalis	-	-	-	-	-	-

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	T					
Chrysosporium sp.	-	-		-	-	-
Cladosporium sp.	•	-	-	-	-	-
Coccidioides immitis	. •	-	-	-	-	-
Cryptococcus	-	-	-	-	-	-
neoformans						
Curvularia sp.	-	-	-	-	•	-
Fusarium sp.	-	-	-	-	-	. -
Geotrichum sp.	-		-	-	•	-
Histoplasma	+	-	-	-	-	-
capsulatum	}					
Mucor sp.	-	-	-	-	-	-
Penicillium sp.		<u>-</u>	-	-	-	-
P. brasiliensis	-	•	-	-	· <u>-</u>	-
Pneumocystis carinii	-	-	-	-	-	-
Pseudallescheria	-	• .	-	-	-	-
boydii						
Rhizopus sp.	-	-	-		-	-
Saccharomyces	•	-			-	-
cerevisiae			·			
Scopulariopsis	-	-	-	-	-	-
brevicaulis		Ī				
Sporothrix schenckii	-	-	-	-	-	-
Trichosporon beigelii	-	-	-	-	-	-

+	Positive
-	Negative after 20 minute wash in 2X SSC

Table 6:

Detection of species specific 28S sequence with probes SEQ ID NO: 15 to SEQ ID

NO: 20

FUNGUS	SEQ ID:	SEQ ID:	SEQ ID:	SEQ ID:	SEQ ID:	SEQ ID:
	15	16	17	18	19	20
Acremonium sp.		-	-	<u>-</u>	-	-
Aspergillus clavatus	-	-	-	7	-	<u>.</u>
Aspergillus flavus	-	-	-		-	-
Aspergillus fumigatus	• .	-	-	-	-	-
Aspergillus glaucus	-	-	-	• .	<u>.</u>	-
Aspergillus nidulans	-	-	-	- .	-	-
Aspergillus niger	-	~	-	-	•	-
Aspergillus ochraceus	-	· •	-	-	-	-
Aspergillus terreus	-	-	-	-	•	-
Aspergillus unguis	-	-	•	•	-	-
Aspergillus ustus	-	-	-	•	-	-
Aspergillus sp.	-	-	· -	-	-	-
Beauvaria sp.	-	-	-	-		•
Bipolaris sp.	-	-	-	-	-	-
Blastomyces	-	•	-	-	-	-
dermatitidis						
Candida albicans	-	•	-	-	-	• 17
Candida glabrata	-	-	-	-	-	-
Candida	-	-		- ·	-	-
guilliermondii			-			
Candida kefyr	+	-	-	-	-	-
Candida krusei	-	+	-	- ·	-	-
Candida lusitaniae	-	-	+	-	-	-
Candida parapsilosis	•		-	+	<u>-</u>	-
Candida tropicalis		-	-	-	+	• ,

					•	
Chrysosporium sp.	•	-			-	
Cladosporium sp.	-	-	-	-	-	-
Coccidioides immitis	-	-	<u> </u>		-	-
Cryptococcus	-	-		-	-	-
neoformans	* .					
Curvularia sp.	-	-	-	-	-	-
Fusarium sp.		· •	-	-	-	
Geotrichum sp.	-	-	-	-		-
Histoplasma	-	-	-	-	•	
capsulatum						•
Mucor sp.	-		-	-	-	•
Penicillium sp.	-	-	-	-	-	· •
P. brasiliensis		-	-	•	· -	-
Pneumocystis carinii	-	-	-	-	-	-
Pseudallescheria	-	-	-	-	-	+
boydii					(
Rhizopus sp.	-	-	-	-	-	-
Saccharomyces	-	-	-	-	-	.=
cerevisiae						
Scopulariopsis	-	-	-	-	-	-
brevicaulis						
Sporothrix schenckii	. •	-	-	-	~	-
Trichosporon beigelii	-	-	-	-	-	-
		L		L		

+ Positive

⁻ Negative after 20 minute wash in 2X SSC

Table 7:
Detection of species specific 28S sequence with probes SEQ ID NO: 21 to SEQ ID NO: 23

FUNGUS	SEQ ID:	SEQ ID:	SEQ ID:
	21	22	23
Acremonium sp.	-	-	-
Aspergillus clavatus	-	-	-
Aspergillus flavus	+	-	
Aspergillus fumigatus	-	-	-
Aspergillus glaucus	-	-	-
Aspergillus nidulans	-	-	-
Aspergillus niger	-	-	-
Aspergillus ochraceus		•	-
Aspergillus terreus	-	-	-
Aspergillus unguis	-	-	-
Aspergillus ustus	-		-
Aspergillus sp.	-	-	-
Beauvaria sp.	-	-	-
Bipolaris sp.	-	-	-
Blastomyces dermatitidis	-	-	-
Candida albicans	-	-	-
Candida glabrata	-	-	-
Candida guilliermondii	-	-	-
Candida kefyr	-	-	-
Candida krusei	-	-	-
Candida lusitaniae	-	-	-
Candida parapsilosis	-	-	-
Candida tropicalis	-	-	-
Chrysosporium sp.	-	-	-
Cladosporium sp.	-	-	-

Coccidioides immitis	- 1	-	<u>-</u>
Cryptococcus neoformans	•	•	-
Curvularia sp.	· -	-	-
Fusarium sp.	-	-	-
Geotrichum sp.	-	-	_
Histoplasma capsulatum	•	•	-
Mucor sp.	-	-	<u>-</u>
Penicillium sp.	•	-	-
Paracoccidioides brasiliensis	-	-	<u>-</u>
Pneumocystis carinii	•	-	-
Pseudallescheria boydii	-	-	_
Rhizopus sp.	·-	•	-
Saccharomyces cerevisiae	-	-	-
Scopulariopsis brevicaulis	-	•	-
Sporothrix schenckii	-	+	+
Trichosporon beigelii	-	-	
			l '

+	Positive	
-	Negative after 20 minute wash in 2X SSC	

Table 8:

Detection of species specific 28S sequence with probes SEQ ID NO: 75 to SEQ ID NO: 76

FUNGUS	SEQ ID NO: 75	SEQ ID NO: 76
Acremonium sp.	-	•
Aspergillus clavatus	-	-
Aspergillus flavus	-	-
Aspergillus fumigatus	-	-
Aspergillus glaucus		-
Aspergillus nidulans	-	• .
Aspergillus niger	-	-
Aspergillus ochraceus		-
Aspergillus terreus	-	-
Aspergillus unguis	-	-
Aspergillus ustus		
Beauvaria sp.	-	-
Bipolaris sp.	-	-
Blastomyces dermatitidis	-	-
Blasotschizomyces capitatus		-
Candida albicans	- 1	
Candida glabrata	-	-
Candida guilliermondii	-	-
Candida krusei	-	-
Candida lusitaniae	-	-
Candida parapsilosis		-
Candida tropicalis	-	-
Chrysosporium sp.	-	
Cladosporium sp.	-	-
Coccidioides immitis	-	-

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Cryptococcus laurentii		•
Cryptococcus neoformans	-	-
var neoformans (sero A)	-	-
var neoformans (sero D)	-	
var gatti (sero B)	-	-
Cryptococcus terreus	-	-
Curvularia sp.	-	-
Filobasidium capsuligenum	-	-
Filobasidium uniguttulatum	- .	-
Fusarium sp.	-	- .
Geotrichum sp.	-	-
Histoplasma capsulatum	-	-
Malbranchea filamentosum	-	•
Mucor sp.	-	-
Paecilomyces sp.		-
Paracoccidioides brasiliensis		
isolate no. 135	+ .	-
isolate no. 262	+	-
isolate no. 265	+	-
isolate no. 927	+	-
isolate no. 9919	+	-
isolate no. 9894	+	-
Prototheca sp.	-	-
Penicillium sp.	-	<u>-</u>
Pneumocystis carinii		
isolate no. Mayo ref. cult.	-	+
isolate no. 56	-	+
isolate no. 62	-	+
isolate no. 69	,-	+
isolate no. 85	-	+
Pseudallesheria boydii	-	-

Rhizopus sp.	-	-
Saccharomyces cerevisiae	-	-
Scopulariopsis brevicaulis	-	-
Scopulariopsis brumptii	-	-
Sporothrix schenckii		
Trichosporon beigelii	-	-

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Table 9:
GenBank search results listing genes from other organisms having 100% identity to probes SEQ ID NO: 3 to SEQ ID NO: 23, SEQ ID NO:75 and SEQ ID NO:76

	PROBE	ORGANISM	GENE	ACCESSION
	SEQ ID	MATCHED	MATCHED*	NUMBER
	NO:		(see note below)	
Aspergillus fumigatus	3	-	-	-
Blastomyces	4	Streptomyces	bleomycin acetyl	L26955
dermatitidis	ļ. 	verticillus	transferase	
	4	Giardia muris	upstream of	X65063, S53320
			rRNA genes	
	4	Aspergillus nidulans	uric acid-	X71807
·			xanthine	
			permease	-
	4	Homo sapiens	T-cell surface	X16996
			glycoprotein	
	4	Homo sapiens	MIC2	M16279,
				M22557,
				J03841, M22556
Candida albicans	5	Candida albicans	28S rRNA	L28817
Coccidioides immitis	6	-	-	•
Cryptococcus	7	Cryptococcus	28S rRNA	L14067,
neoformans		neoformans		L14068,
Cryptococcus	8	Cryptococcus	28S rRNA	L14067,
neoformans		neoformans		L14068, L20964
	8	Escherichia coli	0111 cld	Z17241
Histoplasma	9	.	*	-
capsulatum	*	·		

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Aspergillus glaucus	10	Pseudomonas	cob genes	M62866
·		denitrificans		
Aspergillus niger	11	•	-	•
Aspergillus terreus	12	Human	genome	X17403
·		cytomegalovirus	,	
	12	Homo sapiens	GABA receptor	L08485
Candida glabrata	13	Homo sapiens	Class 1 MHC	X03664,
				X03665
Candida	14	. •	-	-
guilliermondii				
Candida kefyr	15 .	-	-	-
Candida krusei	16	Pseudomonas	penicillin binding	L28837
		syringae	protein	
Candida lusitaniae	17	Chicken	AK1	D00251
	17	Mouse	IL10	M84340
Candida parapsilosis	18	Polytomella agilis	beta-2 tubulin	M33373
	18	Tobacco chloroplast	genome	Z00044, S54304
	18	Aedes aegypti	amylase	L03640
	18	Homo sapiens	chromosome	L14473
			13q14	
Candida tropicalis	19	-	*	-
Pseudallescheria	20	Drosophila	AcTr66B	X71789
boydii		melanogaster		
		Cow	actin 2	D12816
Aspergillus flavus	21	-	-	•
Sporothrix schenckii	22	-	-	-
Sporothrix schenckii	23	Sulfate reducing	FMN binding	D21804
		bacteria	protein	
•	23	Equine herpesvirus 1	genome	M86664
Paracoccidioides	75	European rabbit	BI-1 calcium	X57476
brasiliensis			channel gene	

Pneumocystis carinii	76	rice	heat shock protein	D1775
	76	C. elegens	repetitive DNA	X61259
	76	Synechocystis sp.	genome	D64000
	76	C. elegens	various cosmids	Z81063,
	•			U59749,
				Z69662,
				Z49936,
0		·		U88182,
				U80931,
			·	Z47358,
				Z81142,
				Z79602,
				Z82084,
				Z70269

* Note: As discussed earlier in this document, the presence of sequences similar to probes SEQ ID NO:3 to SEQ ID NO: 23, SEQ ID NO:75 and SEQ ID NO:76 in genes not related to 28S does not have any effect on the specificity or sensitivity of our diagnostic strategy. Our species specific probes are used to analyze 28S DNA that has been previously amplified in a polymerase chain reaction with our probes SEQ ID NO: 1 and SEQ ID NO:2. These two probes will not amplify DNA from any gene other than 28S in column #4 (GENE MATCHED), and therefore no amplified DNA from these non-28S genes will be available for the hybridization of probes SEQ ID NO: 3 to SEQ ID NO: 23, SEQ ID NO:75 and SEQ ID NO:76.

EXAMPLE 2. Use of method in example 1 to test clinical specimens for specific fungal organisms.

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Clinical samples taken from the respiratory and gastrointestinal tract of healthy individuals almost always contain some fungal flora. Most of these fungi are non-

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pathogenic, but may give false positives on traditional immunochemical diagnostic tests for pathogenic fungi.

We obtained 44 clinical specimens from diverse sources ranging from sputum and incision drainage tubes, to intervertebral disc and lung biopsies. Traditional smear and culture results showed that all 44 specimens contained at least 1 type of fungus. In order to test the efficacy of our probes, we extracted DNA from all 44 clinical samples and used probes SEQ ID NO: 1 & 2 in a polymerase chain reaction to amplify fungal 28S sequences present in these samples.

DNA was extracted from all clinical samples by our modification of the technique of Chomczynski and Sacchi which originally described the use of acid guanidinium thiocyanate-phenol-chloroform to preferentially extract RNA from cells and tissues. We replaced room temperature cell lysis by boiling lysis, and acid guanidinium thiocyanate-phenol-chloroform extraction by alkaline phenol-guanidine thiocyanate to preferentially extract DNA from cells. 1.5 ml Sarsted (Newton, North Carolina) polypropylene screw cap tubes with o-ring seals were used for the extractions. 200 ul of specimen was added to 500 ul of GPT reagent (6 M guanidine thiocyanate dissolved in 50 mM tris pH 8.3 mixed with an equal volume of phenol buffered in tris pH 8.0). This was mixed by vortexing and immediately placed in a boiling water bath for 15 minutes. The tubes were spun in a microcentrifuge for 5 seconds and 250 ul of chloroform/iso-amyl alcohol (24:1 by volume) was added and mixed by vortexing. The liquid phases were separated by centrifugation for 10 minutes and 450 ul of aqueous (upper) phase was transferred to a fresh tube. The aqueous phase was mixed with 500 ul of 100% isopropanol and placed at -20°C for at least 1 hour. At the end of this period the tubes were centrifuged for 15 minutes and the supernatant removed without disturbing the nucleic acid pellet. The pellet was washed with 500 ul of ice-cold 70% ethanol to remove traces of GPT reagent by gently inverting 2 times and then centrifuged for 5 minutes. The ethanol was removed and the pellet dried in a speed vac for 10 minutes. The pellet was resuspended in 25 ul of sterile deionized water and 5 ul was used in a 50 ul PCR amplification. The PCR was carried out as a hot-start reaction using the thermal cycling conditions for probes SEQ ID NO: 1 and SEQ ID NO: 2 described in example 1. Gel electrophoresis

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showed that probes SEQ ID NO: 1 and SEQ ID NO: 2 successfully amplified DNA from all 44 specimens.

The amplified DNA from each specimen was transferred to a positively charged polysulphone based membrane. We radioactively labeled our species specific probes SEQ ID NO: 3, SEQ ID NO: 5, SEQ ID NO: 6 and SEQ ID NO: 7, and sequentially probed the membrane to test for the presence of 28S rDNA from Aspergillus fumigatus, Candida albicans, Coccidioides immitis and Cryptococcus neoformans respectively. Membrane blocking, probe hybridization and washes were done exactly as described in Example 1. The results are shown in Table 10.

No false positives were observed, indicating a specificity of 100% for these 4 probes in the clinical specimens tested. 10 out of 12 culture positive samples for *Aspergillus fumigatus*, and 11 out of 13 samples of *Candida albicans* were identified, indicating a detection sensitivity of about 85% for these two probes. Additionally, two out of two *Coccidioides immitis* and two out of two *Cryptococcus neoformans* were correctly identified (detection sensitivity of 100%). As seen by these results, the probes described in this invention can be used on a diverse variety of clinical specimens with excellent efficacy.

Table 10.

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Detection of Aspergillus fumigatus, Candida albicans, Coccidioides immitis and Cryptococcus neoformans in clinical specimens using species specific probes.

Specimen type	Smear and	PCR with	SEQ ID:	SEQ ID:	SEQ ID:	SEQ ID:
	culture	SEQ ID: 1,	3	5	6	7
·	results	2				
U035 sputum	A. flavus	+	-	•	-	-
U069 pleura	A. fumigatus	+	+	-	-	-
U070 bronchial wash	A. flavus	+	*	-	-	• .

M019 bronchial	A. fumigatus	+	+	T -	-	T -
wash		. ,				
M020 sputum	mixed fungal	+	-	+	•	-
	flora			•		
X35254 sputum	C. albicans	+	-	+	-	-
M20910 sputum	A. fumigatus	+	+	-	-	-
M055 sputum	C. albicans	+	-	+	-	-
M056 abdominal	mixed fungal	+		•	-	-
	flora					
M057 drainage	C. albicans	+	- .	(-)	:	-
tube			. •			
M059 ind. sputum	C. albicans	+	-	+	-	•
M060 ind. sputum	mixed fungal	+	-	-	-	-
·	flora					
M083 bronchial	C. albicans	+	-	+	-	<u>-</u>
wash						
M084 sputum	A. fumigatus	+	(-)	-	-	<u>-</u>
M085 throat	C. albicans	+	-	(-)	-	-
A001 sputum	A. fumigatus	+	(-)	-	-	- .
A002 leg	Blastomyces	+	-	-	-	-
A003 leg	Blastomyces	+	-	-	-	-
A005 disc	A. fumigatus	+	+		-	-
A037 disc	A. fumigatus	+	+		-	-
A039 trachea	C. albicans	+	-	+	-	-
A040 trachea	C. albicans	+	•	+	-	-
A102 empyema	A. fumigatus	+	+	_	-	-
Y004 sputum	C. albicans	+	•	+	-	-
Y016 induced	Coccidioides	+	•	-	+	-
sputum					·	
Y028 sputum	Coccidioides	+	•	-	+	-
J003 chest	Aspergillus sp.	+	-	-	-	-

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J045 bronchial	C. albicans	+		+	T	<u> </u>
`	C. aibicaris		-	T	-	•
wash	.]					
J046 ethmoid	yeast	+		-	-	-
J047 chest	A. fumigatus	+	+	-	-	-
J048 sputum	C. albicans	+	-	+ ,	-	-
J073 lung	Aspergillus sp.	+	-	•	-	-
J074 lung	A. fumigatus	+	+	-		-
U017 lip	A. fumigatus	+	+	-	. ~	-
U033 sputum	mixed fungal	+	-	-	-	-
	flora					
U071 sputum	C. albicans	+	-	+	-	_
U072 BA lavage	Sporothrix	+	-	•	-	-
U073 knee	Histoplasma	+	-	-	-	-
U074 mandible	Cryptococcus	+	-	- ,	-	+
U075 CSF	Cryptococcus	+	-	-	-	+
U076 knee	Histoplasma	+	-	-	-	-
U077 soft tissue	Histoplasma	+	-	- .	-	-
U051 buccal	A. fumigatus	+	+	-	-	-
Y055 sputum	mixed fungal	+	-	**	-	-
	flora					
+ Positive -	Negative (-) I	Missed				L

EXAMPLE 3. DNA sequence based identification of unknown fungal organisms.

Another utility of our probes is in the rapid DNA sequence based identification of a pure culture of fungus. Probes SEQ ID NO: 1 and SEQ ID NO: 2 are used in a polymerase chain reaction to amplify 28S rDNA from an unknown fungus. Probes SEQ ID NO: 1 or SEQ ID NO: 2 are then used as sequencing primers to obtain DNA sequence from this amplified 28S DNA belonging to the unknown fungus. This DNA sequence is compared to the fungal 28S DNA sequences in our database, and a sequence match at, or overlapping any one of the probe sequences in SEQ ID NO: 3 to

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SEQ ID NO: 78 will confirm the identity of the fungus. This technique cannot be used directly on clinical samples, as these usually contain DNA from more than one fungus, and the DNA sequence generated will consist of overlapping sequences of several organisms. This technique has utility in rapidly and reliably identifying colonies of a single fungus on culture plates, clinical specimens, food, pharmaceutical, environmental or other samples containing only one species of fungus.

EXAMPLE 4. Capture and identification of target DNA or RNA

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All primers and probes described in this invention disclosure may be labeled with any detectable reporter or signal moiety including, but not limited to radioisotopes, enzymes, antigens, antibodies, chemiluminescent reagents and fluorescent chemicals. Additionally, these probes may be modified without changing the substance of their purpose by terminal addition of nucleotides designed to incorporate restriction sites or other useful sequences. These probes may also be modified by the addition of a capture moiety (including, but not limited to paramagnetic particles, biotin, fluorescein, dioxigenin, antigens, antibodies) or attached to the walls of microtiter trays to assist in the solid phase capture and purification of these probes and any DNA or RNA hybridized to these probes. Fluorescein may be used as a signal moiety as well as a capture moiety, the latter by interacting with an anti-fluorescein antibody.

A typical utility of these modifications would be as follows. Primers SEQ ID NO: 1 and SEQ ID NO: 2 would be utilized to amplify 28S rDNA from a sample, if present, as described previously. Primers would be modified so as to contain a biotin moiety at their 5' ends. A streptavidin solid phase, such as a paramagnetic particle, would be used to separate PCR products, if present, from the reaction mixture. The amplified target may be subsequently hybridized to a third probe ((SEQ ID NO: 3) to (SEQ ID NO: 78) or their complements) attached to a detectable moiety to determine which species of fungus is present in the given sample. Multiple probes, each labeled with a different detectable moiety may be used at one time to analyze the amplified target.

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Alternatively, Primers SEQ ID NO: 1 and SEQ ID NO: 2 would be utilized to amplify 28S rDNA from a sample, if present, as above. In a separate reaction, individually, either SEQ ID NO: 1 or SEQ ID NO: 2 would be modified by attachment to a solid phase capture moiety, such as a paramagnetic particle, and SEQ ID NO: 3 to SEQ ID NO: 78 (or their complements) would be modified by addition of a detectable moiety. Alternately, in the amplicon, any sequences delimited by SEQ ID NO: 1 and SEQ ID NO: 2, including but not limited to SEQ ID NO: 3 to SEQ ID NO: 78, may be used in the design of a capture probe. One of the probes attached to a solid phase (SEQ ID NO: 1 and SEQ ID NO: 2) or any other appropriately designed sequences and one of the probes modified by attachment to a detectable moiety (SEQ ID NO: 3 to SEQ ID NO: 78 or their complements) would be hybridized together, in solution, to products of the PCR, if they had been generated. The hybrids, if present, would be captured from the solution, and analyzed by a method appropriate to the detection moiety. Detection of the hybridized probe would indicate which species of fungus was present in the given sample. Multiple probes, each labeled with a different detectable moiety may be used at one time to analyze the amplified target.

EXAMPLE 5. Species-specific amplification of fungal DNA

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Another utility of the probes described in this invention is their usage as primers in the direct detection of a specific fungal species by virtue of a nucleic acid amplification reaction. In this embodiment, one primer is a universal one, such as (SEQ ID NO:1) or (SEQ ID NO:2), and the other is a species-specific primer selected from the group consisting of (SEQ ID NO:3) to (SEQ ID NO: 23), (SEQ ID NO:75) and (SEQ ID NO:76) or the complements thereof. One variation of this approach is the substitution of (SEQ ID NO:1) or (SEQ ID NO:2) with any functional sequence located in proximity to the species-specific primer. Another variation of this approach is the selection of any appropriate species specific primer pair from SEQ ID NO: 24 to SEQ ID NO: 74, SEQ ID NO:77 and SEQ ID NO:78.

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SEQUENCE LISTING

(1) GENERAL INFORMATION:

- (i) APPLICANT:
 - (A) Sandhu, Gurpreet S.
 - (B) Kline, Bruce C.
- (ii) TITLE OF INVENTION:

Nucleic Acid Probes for the Detection and Identification of Fungi

- (iii) NÚMBER OF SEQUENCES: 80
- (iv) CORRESPONDENCE ADDRESS:
 - (A) ADDRESSEE: Chiron Diagnostics Corporation
 - (B) STREET: 63 North Street
 - (C) CITY: Medfield
 - (D) STATE: Massachusetts
 - (E) COUNTRY: USA
 - (F) ZIP: 02052
 - (v) COMPUTER READABLE FORM:
 - (A) MEDIUM TYPE: Diskette 3.5 inch, 1.44 Mb storage
 - (B) COMPUTER: IBM PS/2
 - (C) OPERATING SYSTEM: MS-DOS 6.2
 - (D) SOFTWARE: Word 6.0
 - (vi) CURRENT APPLICATION DATA:
 - (A) APPLICATION NUMBER:
 - (B) FILING DATE:
 - (C) CLASSIFICATION:

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(vii) PRIOR APPLICATION DATA:

- (A) APPLICATION NUMBER: 08/373,127
- (B) FILING DATE: 13-JAN-1995
- (C) CLASSIFICATION

(vii) PRIOR APPLICATION DATA:

- (A) APPLICATION NUMBER: 08/435,684
- (B) FILING DATE: 05-MAY-1995
- (C) CLASSIFICATION

(viii) ATTORNEY INFORMATION:

- (A) NAME: Morgenstern, Arthur S.
- (B) REGISTRATION NUMBER: 28,244
- (C) DOCKET NUMBER: CCD-180CIPII

(ix) TELECOMMUNICATION INFORMATION:

- (A) TELEPHONE: 508 359-3836
- (B) TELEFAX: 508 359-3885

(2) INFORMATION FOR SEQ ID NO 1:

- (i) SEQUENCE CHARACTERISTICS:
 - (A) LENGTH: 20
 - (B) TYPE: nucleic acid
 - (C) STRANDEDNESS: single
 - (D) TOPOLOGY: linear
- (ii) MOLECULE TYPE: Nucleic acid probe for fungal organisms

(iii) HYPOTHETICAL: No

- (iv) ANTISENSE: No
- (v) SEQUENCE DESCRIPTION: SEQ ID NO: 1:
- GTGAAATTGT TGAAAGGGAA

- (3) INFORMATION FOR SEQ ID NO 2:
 - (i) SEQUENCE CHARACTERISTICS:
 - (A) LENGTH: 18
 - (B) TYPE: nucleic acid
 - (C) STRANDEDNESS: single
 - (D) TOPOLOGY: linear
 - (ii) MOLECULE TYPE: Nucleic acid probe for fungal organisms
- (iii) HYPOTHETICAL: No

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- (iv) ANTISENSE: No
- 20 (v) SEQUENCE DESCRIPTION: SEQ ID NO: 2:
 - GACTCCTTGG TCCGTGTT

- (4) INFORMATION FOR SEQ ID NO 3:
 - (i) SEQUENCE CHARACTERISTICS:
 - (A) LENGTH: 14
 - (B) TYPE: nucleic acid
 - (C) STRANDEDNESS: single
- (D) TOPOLOGY: linear
 - (ii) MOLECULE TYPE: Nucleic acid probe for Aspergillus fumigatus

	(iii) HYPOTHETICAL: No
	(iv) ANTISENSE: No
5	(v) SEQUENCE DESCRIPTION: SEQ ID NO: 3:
	CTCGGAATGT ATCA 14
10	(5) INFORMATION FOR SEQ ID NO 4:
	(i) SEQUENCE CHARACTERISTICS:
	(A) LENGTH: 13
	(B) TYPE: nucleic acid
15	(C) STRANDEDNESS: single
	(D) TOPOLOGY: linear
	(ii) MOLECULE TYPE: Nucleic acid probe for Blastomyces dermatitidis
20	(iii) HYPOTHETICAL: No
	(iv) ANTISENSE: No
25	(v) SEQUENCE DESCRIPTION: SEQ ID NO: 4:
25	ACTCCCCCAC GGG 13
	(6) INFORMATION FOR SEQ ID NO 5:
30	(i) SEQUENCE CHARACTERISTICS:
	(A) LENGTH: 14
	(B) TYPE: nucleic acid

	(C) STRANDEDNESS: single (D) TOPOLOGY: linear
	(ii) MOLECULE TYPE: Nucleic acid probe for Candida albicans
5	(iii) HYPOTHETICAL: No
	(iv) ANTISENSE: No
10	(v) SEQUENCE DESCRIPTION: SEQ ID NO: 5:
	CCTCTGACGA TGCT 14
	(7) INFORMATION FOR SEQ ID NO 6:
15	(i) SEQUENCE CHARACTERISTICS: (A) LENGTH: 14
	(B) TYPE: nucleic acid
20	(C) STRANDEDNESS: single (D) TOPOLOGY: linear
	(ii) MOLECULE TYPE: Nucleic acid probe for Coccidioides immitis
	(iii) HYPOTHETICAL: No
25	(iv) ANTISENSE: No
	(v) SEQUENCE DESCRIPTION: SEQ ID NO: 6:
30	TCTGGCGGTT GGTT 14
	(8) INFORMATION FOR SEQ ID NO 7:
	(i) SEQUENCE CHARACTERISTICS:

(A) LENGTH: 14

	(B) TYPE: nucleic acid
	(C) STRANDEDNESS: single
٠	(D) TOPOLOGY: linear
5	
	(ii) MOLECULE TYPE: Nucleic acid probe for Cryptococcus neoformans
	(iii) HYPOTHETICAL: No
0	(iv) ANTISENSE: No
	(v) SEQUENCE DESCRIPTION: SEQ ID NO: 7:
5	CTCCTGTCGC ATAC 14
	(9) INFORMATION FOR SEQ ID NO 8:
	(i) SEQUENCE CHARACTERISTICS:
	(A) LENGTH: 14
0	(B) TYPE: nucleic acid
	(C) STRANDEDNESS: single
	(D) TOPOLOGY: linear
5	(ii) MOLECULE TYPE: Nucleic acid probe for Cryptococcus neoformans
	(iii) HYPOTHETICAL: No
	(iv) ANTISENSE: No
	(v) SEQUENCE DESCRIPTION: SEQ ID NO: 8:
0	AGTTCTGATC GGTG 14

(10) INFORMATION FOR SEQ ID NO 9:

- (i) SEQUENCE CHARACTERISTICS:
 - (A) LENGTH: 14
 - (B) TYPE: nucleic acid
 - (C) STRANDEDNESS: single
 - (D) TOPOLOGY: linear
- (ii) MOLECULE TYPE: Nucleic acid probe for Histoplasma capsulatum
- (iii) HYPOTHETICAL: No
 - (iv) ANTISENSE: No
- (v) SEQUENCE DESCRIPTION: SEQ ID NO: 9:

CAATCCCCG CGGC

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(11) INFORMATION FOR SEQ ID NO 10:

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- (i) SEQUENCE CHARACTERISTICS:
 - (A) LENGTH: 14
 - (B) TYPE: nucleic acid
 - (C) STRANDEDNESS: single
 - (D) TOPOLOGY: linear
- (ii) MOLECULE TYPE: Nucleic acid probe for Aspergillus glaucus
- (iii) HYPOTHETICAL: No
- (iv) ANTISENSE: No

(v) SEQUENCE DESCRIPTION: SEQ ID NO: 10:

GTGTCATGCG GCCA

14

- (12) INFORMATION FOR SEQ ID NO 11:
 - (i) SEQUENCE CHARACTERISTICS:
 - (A) LENGTH: 14
 - (B) TYPE: nucleic acid
 - (C) STRANDEDNESS: single
 - (D) TOPOLOGY: linear
 - (ii) MOLECULE TYPE: Nucleic acid probe for Aspergillus niger
- 15 (iii) HYPOTHETICAL: No
 - (iv) ANTISENSE: No
 - (v) SEQUENCE DESCRIPTION: SEQ ID NO: 11:

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CCCTGGAATG TAGT

14

- (13) INFORMATION FOR SEQ ID NO 12:
 - (i) SEQUENCE CHARACTERISTICS:

(A) LENGTH: 14

(B) TYPE: nucleic acid

(C) STRANDEDNESS: single

(D) TOPOLOGY: linear

- (ii) MOLECULE TYPE: Nucleic acid probe for Aspergillus terreus
- (iii) HYPOTHETICAL: No

	(iv) ANTISENSE: No
	(v) SEQUENCE DESCRIPTION: SEQ ID NO: 12:
5	GCTTCGGCCC GGTG 14
	(14) INFORMATION FOR SEQ ID NO 13:
10	(i) SEQUENCE CHARACTERISTICS:
	(A) LENGTH: 14
	(B) TYPE: nucleic acid
	(C) STRANDEDNESS: single
	(D) TOPOLOGY: linear
15	(ii) MOLECULE TYPE: Nucleic acid probe for Candida glabrata
	(iii) HYPOTHETICAL: No
	(iv) ANTISENSE: No
20	(v) SEQUENCE DESCRIPTION: SEQ ID NO: 13:
	CTTGGGACTC TCGC 14
25	(15) INFORMATION FOR SEQ ID NO 14:
	(i) SEQUENCE CHARACTERISTICS:
	(A) LENGTH: 14
	(B) TYPE: nucleic acid
30	(C) STRANDEDNESS: single
	(D) TOPOLOGY: linear

	(ii) MOLECULE TYPE: Nucleic acid probe for Candida guilliermondii
	(iii) HYPOTHETICAL: No
5	(iv) ANTISENSE: No
	(v) SEQUENCE DESCRIPTION: SEQ ID NO: 14:
•	ATATTTTGTG AGCC 14
0	(16) INFORMATION FOR SEQ ID NO 15:
	(i) SEQUENCE CHARACTERISTICS:
	(A) LENGTH: 14
5	(B) TYPE: nucleic acid
	(C) STRANDEDNESS: single
	(D) TOPOLOGY: linear
0	(ii) MOLECULE TYPE: Nucleic acid probe for Candida kefyr
U	(iii) HYPOTHETICAL: No
	(iv) ANTISENSE: No
5	(v) SEQUENCE DESCRIPTION: SEQ ID NO: 15:
	TTCGGCTTTC GCTG 14
0	(17) INFORMATION FOR SEQ ID NO 16:
	(i) SEQUENCE CHARACTERISTICS:

(A) LENGTH: 14

	(B) TYPE: nucleic acid
	(C) STRANDEDNESS: single
	(D) TOPOLOGY: linear
5	(ii) MOLECULE TYPE: Nucleic acid probe for Candida krusei
	(iii) HYPOTHETICAL: No
	(iv) ANTISENSE: No
10	(v) SEQUENCE DESCRIPTION: SEQ ID NO: 16:
	GGGATTGCGC ACCG 14
	(18) INFORMATION FOR SEQ ID NO 17:
15	
	(i) SEQUENCE CHARACTERISTICS:
	(A) LENGTH: 14
	(B) TYPE: nucleic acid
	(C) STRANDEDNESS: single
20	(D) TOPOLOGY: linear
	(ii) MOLECULE TYPE: Nucleic acid probe for Candida lusitaniae
25	(iii) HYPOTHETICAL: No
23	(iv) ANTISENSE: No
	(v) SEQUENCE DESCRIPTION: SEQ ID NO: 17:
30	GCCTCCATCC CTTT 14

(19) INFORMATION FOR SEQ ID NO 18:

	(A) LENGTH: 14
	(B) TYPE: nucleic acid
5	(C) STRANDEDNESS: single
	(D) TOPOLOGY: linear
	(ii) MOLECULE TYPE: Nucleic acid probe for Candida parapsilosis
10 .	(iii) HYPOTHETICAL: No
	(iv) ANTISENSE: No
	(v) SEQUENCE DESCRIPTION: SEQ ID NO: 18:
15	ATAAGTGCAA AGAA 14
	(20) INFORMATION FOR SEQ ID NO 19:
	(i) SEQUENCE CHARACTERISTICS:
20	(A) LENGTH: 14
	(B) TYPE: nucleic acid
	(C) STRANDEDNESS: single
	(D) TOPOLOGY: linear
25	(ii) MOLECULE TYPE: Nucleic acid probe for Candida tropicalis
	(iii) HYPOTHETICAL: No
20	(iv) ANTISENSE: No
30	(v) SEQUENCE DESCRIPTION: SEQ ID NO: 19:

(i) SEQUENCE CHARACTERISTICS:

- (21) INFORMATION FOR SEQ ID NO 20:
 - (i) SEQUENCE CHARACTERISTICS:
 - (A) LENGTH: 14
 - (B) TYPE: nucleic acid
 - (C) STRANDEDNESS: single
 - (D) TOPOLOGY: linear
 - (ii) MOLECULE TYPE: Nucleic acid probe for Pseudallescheria boydii
 - (iii) HYPOTHETICAL: No
 - (iv) ANTISENSE: No
 - (v) SEQUENCE DESCRIPTION: SEQ ID NO: 20:

GCGATGGGAA TGTG

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- 20 (22) INFORMATION FOR SEQ ID NO 21:
 - (i) SEQUENCE CHARACTERISTICS:
 - (A) LENGTH: 14
 - (B) TYPE: nucleic acid
 - (C) STRANDEDNESS: single
 - (D) TOPOLOGY: linear
 - (ii) MOLECULE TYPE: Nucleic acid probe for Aspergillus flavus
- (iii) HYPOTHETICAL: No
 - (iv) ANTISENSE: No

(v) SEQUENCE DESCRIPTION: SEQ ID NO: 21:

AGACTCGCCT CCAG

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- (23) INFORMATION FOR SEQ ID NO 22:
 - (i) SEQUENCE CHARACTERISTICS:
 - (A) LENGTH: 14
 - (B) TYPE: nucleic acid.
 - (C) STRANDEDNESS: single
 - (D) TOPOLOGY: linear
 - (ii) MOLECULE TYPE: Nucleic acid probe for Sporothrix schenckii
- (iii) HYPOTHETICAL: No
 - (iv) ANTISENSE: No
 - (v) SEQUENCE DESCRIPTION: SEQ ID NO: 22:

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CGGACCACCC GGCG

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- (24) INFORMATION FOR SEQ ID NO 23:
 - (i) SEQUENCE CHARACTERISTICS:
 - (A) LENGTH: 14
 - (B) TYPE: nucleic acid
 - (C) STRANDEDNESS: single
 - (D) TOPOLOGY: linear

(ii) MOLECULE TYPE: Nucleic acid probe for Sporothrix schenckii

- (iii) HYPOTHETICAL: No
- (iv) ANTISENSE: No
- (v) SEQUENCE DESCRIPTION: SEQ ID NO: 23:
- CGGCGGCATG CCCC

- (25) INFORMATION FOR SEQ ID NO 24:
 - (i) SEQUENCE CHARACTERISTICS:
 - (A) LENGTH: 208
 - (B) TYPE: nucleic acid
 - (C) STRANDEDNESS: single
 - (D) TOPOLOGY: linear
 - (ii) MOLECULE TYPE: Acremonium species specific region of 28S gene.
 - (iii) HYPOTHETICAL: No
 - (iv) ANTISENSE: No

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(v) SEQUENCE DESCRIPTION: SEQ ID NO: 24:

GACCAGACTT GGGCTCGGTG AATCATCCGG CGTTCTCGCC
25 GGTGCACTTT 50

GCCGTCCCAG GCCAGCATCA GTTCGCGCCG GGGGATAAAG GTTTCGGGAA 100

TGTAGCTCCT TCGGGAGTGT TATAGCCCGT TGCGTAATAC
CCTGGCGTGG 150

ACTGAGGTCC GCGCTCTGCA AGGATGCTGG CGTAATGGTC
ATCAGTGACC 200

CGTCTTGA

(26) INFORMATION FOR SEQ ID NO 25:

- (i) SEQUENCE CHARACTERISTICS:
 - (A) LENGTH: 212
 - (B) TYPE: nucleic acid
 - (C) STRANDEDNESS: single
 - (D) TOPOLOGY: linear

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- (ii) MOLECULE TYPE: Aspergillus clavatus specific region of 28S gene.
- (iii) HYPOTHETICAL: No
- (iv) ANTISENSE: No
- (v) SEQUENCE DESCRIPTION: SEQ ID NO: 25:
- 20 GACCAGACTC GCTCGCGGGG TTCAGCCGGC ATTCGTGCCG
 GTGTACTTCC 50

CCGTGGGCGG GCCAGCGTCG GTTTGGGCGG CCGGTCAAAG GCCTCCGGAA 100

TGTATCACCT CTCGGGGTGT CTTATAGCCG GGGGTGCAAT
GCGGCCTGCC 150

TGGACCGAGG AACGCGCTTC GGCTCGGACG CTGGCGTAAT GGTCGTAAAT 200

GACCCGTCTT GA

212

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(27) INFORMATION FOR SEQ ID NO 26:

- (i) SEQUENCE CHARACTERISTICS:
 - (A) LENGTH: 212
 - (B) TYPE: nucleic acid
 - (C) STRANDEDNESS: single
 - (D) TOPOLOGY: linear
- (ii) MOLECULE TYPE: Aspergillus flavus specific region of 28S gene.
- (iii) HYPOTHETICAL: No
- (iv) ANTISENSE: No
- (v) SEQUENCE DESCRIPTION: SEQ ID NO: 26:

GACCAGACTC GCCTCCAGGG TTCAGCCGGC ATTCGTGCCG

GTGTACTTCC 50

CTGGGGGCGG GCCAGCGTCG GTTTGGGCGG CCGGTCAAAG GCTCCCGGAA 100

TGTAGTGCCC TYCGGGGCAC CTTATAGCCG GGAGTGCAAT GCGGCCAGCC 150

TGGACCGAGG AACGCGCTTC GGCACGGACG CTGGCATAAT
GGTCGYAAAC 200

GACCCGTCTT GA

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212

(28) INFORMATION FOR SEQ ID NO 27:

(i) SEQUENCE CHARACTERISTICS:

- (A) LENGTH: 212
- (B) TYPE: nucleic acid
- (C) STRANDEDNESS: single
- (D) TOPOLOGY: linear
- (ii) MOLECULE TYPE: Aspergillus fumigatus specific region of 28S gene.

- (iii) HYPOTHETICAL: No
- (iv) ANTISENSE: No
- (v) SEQUENCE DESCRIPTION: SEQ ID NO: 27:

GACCAGACTC GCCCGCGGGG TTCAGCCGGC ATTCGTGCCG

GTGTACTTCC 50

CCGTGGCGG GCCAGCGTCG GTTTGGGCGG CCGGTCAAAG GCCCTCGGAA 100

TGTATCACCT CTCGGGGTGT CTTATAGCCG AGGGTGCAAT GCGGCCTGCC 150

TGGACCGAGG AACGCGCTTC GGCTCGGACG CTGGCGTAAT
GGTCGTAAAT 200

GACCCGTCTT GA

212

- 20 (29) INFORMATION FOR SEQ ID NO 28:
 - (i) SEQUENCE CHARACTERISTICS:
 - (A) LENGTH: 212
 - (B) TYPE: nucleic acid
 - (C) STRANDEDNESS: single
 - (D) TOPOLOGY: linear
 - (ii) MOLECULE TYPE: Aspergillus glaucus specific region of 28S gene.
 - (iii) HYPOTHETICAL: No

(iv) ANTISENSE: No

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(v) SEQUENCE DESCRIPTION: SEQ ID NO: 28:

GACCAGACTC GCTTCCGGGG TTCAGCCGGC TTTCGGGCCG
GTGTACTTCC 50

CCGGGGCGG GCCAGCGTCG GTTTGGGCGG CCGGTCAAAG
GCCCCTGGAA 100

TGTAACGCCT CTCGGGGCGC CTTATAGCCA GGGGTGTCAT GCGGCCAGCC 150

TGGACCGAGG AACGCGCTTC GGCACGGACG CTGGCATAAT
GGTCGTAAAC 200

GACCCGTCTT GA

212

(30) INFORMATION FOR SEQ ID NO 29:

- (i) SEQUENCE CHARACTERISTICS:
 - (A) LENGTH: 213
 - (B) TYPE: nucleic acid
 - (C) STRANDEDNESS: single
 - (D) TOPOLOGY: linear
- (ii) MOLECULE TYPE: Aspergillus nidulans specific region of 28S gene.
- 25 (iii) HYPOTHETICAL: No
 - (iv) ANTISENSE: No
 - (v) SEQUENCE DESCRIPTION: SEQ ID NO: 29:

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GACCAGACTC GGCCCCGGGG TTCARCCAGC ACTCGTGCTG
GTGTACTTCC 50

CCGGGGCGG GCCAGCGTCG GTTTGGGCGG CCGGTCAAAG GCCCCAGGAA 100

TGTATCGCCC TCCGGGGTTG TCTTATAGCC TGGGGTGCAA
TGCGGCCAGC 150

CCGGACCGAG GAACGCGCTT CGGCACGGAC GCTGGCGTAA
TGGTCGCAAA 200

CGACCCGTCT TGA

213

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(31) INFORMATION FOR SEQ ID NO 30:

- (i) SEQUENCE CHARACTERISTICS:
 - (A) LENGTH: 212
 - (B) TYPE: nucleic acid
 - (C) STRANDEDNESS: single
 - (D) TOPOLOGY: linear
- (ii) MOLECULE TYPE: Aspergillus niger specific region of 28S gene.
- (iii) HYPOTHETICAL: No
- (iv) ANTISENSE: No

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(v) SEQUENCE DESCRIPTION: SEQ ID NO: 30:

GACCAGACTC GCCCGCGGGG TTCAGCCGGC ATTCGTGCCG
GTGTACTTCC 50

CCGTGGGCGG GCCAGCGTCG GTTTGGGCGG CCGGTCAAAG
GCCCCTGGAA 100

WO 98/55649 PCT/IB98/00865

TGTAGTRCCC TCCGGGGYAC CTTATAGCCA GGGGTGCAAT GCGGCCAGCC 150

TGGACCGAGG AACGCGCTTC GGCACGGACG CTGGCATAAT GGTCGTAAAC 200

GACCCGTCTT GA

212

(32) INFORMATION FOR SEQ ID NO 31:

- (i) SEQUENCE CHARACTERISTICS:
 - (A) LENGTH: 212
 - (B) TYPE: nucleic acid
 - (C) STRANDEDNESS: single
 - (D) TOPOLOGY: linear

(ii) MOLECULE TYPE: Aspergillus ochraceus specific region of 28S gene.

- (iii) HYPOTHETICAL: No
- (iv) ANTISENSE: No

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(v) SEQUENCE DESCRIPTION: SEQ ID NO: 31:

GACCAGACTC GCCCGCGGGG TTCAGCCGGC ATTCGTGCCG
25 GTGTACTTCC 50

CCGCGGCGG GCCAGCGTCG GTTTGGGCGG CCGGTCAAAG GCCCCCGGAA 100

TGTAGCACCC TTCGGGGTGC CTTATAGCCG GGGGTGCAAT
GCGGCCAGCC 150

TGGACCGAGG AACGCGCTTC GGCACGGACG CTGGCATAAT
GGTCGTAAAC 200

GACCCGTCTT GA

(33) INFORMATION FOR SEQ ID NO 32:

- (i) SEQUENCE CHARACTERISTICS:
 - (A) LENGTH: 212
 - (B) TYPE: nucleic acid
 - (C) STRANDEDNESS: single.
 - (D) TOPOLOGY: linear

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- (ii) MOLECULE TYPE: Aspergillus terreus specific region of 28S gene.
- (iii) HYPOTHETICAL: No
- (iv) ANTISENSE: No
- (v) SEQUENCE DESCRIPTION: SEQ ID NO: 32:

AACCAGACTC GCTCGCGGGG TTCAGCCGGG CTTCGGCCCG
20 GTGTACTTCC 50

CCGCGGCGG GCCAGCGTCG GTTTGGGCGG CCGGTCAAAG GCCTCCGGAA 100

TGTAGCGCCC TTCGGGGCGC CTTATAGCCG GGGGTGCAAT GCGGCCAGCC 150

TGGACCGAGG AACGCGCTTC GGCACGGACG CTGGCATAAT
GGTTGTAAAC 200

GACCCGTCTT GA

- 30 (34) INFORMATION FOR SEQ ID NO 33:
 - (i) SEQUENCE CHARACTERISTICS:

(A) LENGTH: 213

(B) TYPE: nucleic acid

(C) STRANDEDNESS: single

(D) TOPOLOGY: linear

(ii) MOLECULE TYPE: Aspergillus unguis specific region of 28S gene.

(iii) HYPOTHETICAL: No

(iv) ANTISENSE: No

(v) SEQUENCE DESCRIPTION: SEQ ID NO: 33:

GACCAGACTC GGCCTCGGGG TTCAGCCAGC ACTCGTGCTG
GTGTACTTCC 50

CCGGGGCGG GCCAGCGTCG GTTTGGGCGG CCGGTCAAAG GCCCCAGGAA 100

TGTATCACCC TCCGGGGTTG TCTTATAGCC TGGGGTGCAA
TGCGGCCAGC 50

CTGGACCGAG GAACGCGCTT CGGCACGGAC GCTGGCATAA TGGTTGCAAA 200

CGACCCGTCT TGA 213

(35) INFORMATION FOR SEQ ID NO 34:

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- (i) SEQUENCE CHARACTERISTICS:
 - (A) LENGTH: 212
 - (B) TYPE: nucleic acid
 - (C) STRANDEDNESS: single
 - (D) TOPOLOGY: linear

- (ii) MOLECULE TYPE: Aspergillus ustus specific region of 28S gene.
- (iii) HYPOTHETICAL: No
- (iv) ANTISENSE: No
- (v) SEQUENCE DESCRIPTION: SEQ ID NO: 34:
- GACCAGACTC GGCCCCGGGG TTCAGCCAGC ACTCGTGCTG
 GTGTACTTCC 50

CCGGGGCGG GCCAGCGTCG GTTTGGGCGG CCGGTCAAAG GCCCCAGGAA 100

TGTGTCGCCC TCCGGGGCGT CTTATAGCCT GGGGTGCAAT GCGGCCAGCC 150

CGGACCGAGG AACGCGCTTC GGCACGGACG CTGGCGTAAT
GGTCGCAAAC 200

GACCCGTCTT GA

212

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- (36) INFORMATION FOR SEQ ID NO 35:
 - (i) SEQUENCE CHARACTERISTICS:
 - (A) LENGTH: 208
 - (B) TYPE: nucleic acid
 - (C) STRANDEDNESS: single
 - (D) TOPOLOGY: linear
 - (ii) MOLECULE TYPE: Beauveria species specific region of 28S gene.
 - (iii) HYPOTHETICAL: No

(v) SEQUENCE DESCRIPTION: SEQ ID NO: 35:

GACCAGACTT GGGCTTGGTT GATCATCCGG GGTTCTCCCC
GGTGCACTCT 50

TCCGGCCCAG GCCAGCATCA GTTCGCCCTG GGGGACAAAG GCTTCGGGAA 100

CGTGGCTCTC TCCGGGGAGT GTTATAGCCC GTTGCGTAAT
ACCCTGTGGC 150

GGACTGAGGT TCGCGCATTC GCAAGGATGC TGGCGTAATG
GTCATCAGTG 200

ACCCGTCT

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208

(37) INFORMATION FOR SEQ ID NO 36:

- (i) SEQUENCE CHARACTERISTICS:
 - (A) LENGTH: 213
 - (B) TYPE: nucleic acid
 - (C) STRANDEDNESS: single
 - (D) TOPOLOGY: linear

(ii) MOLECULE TYPE: Bipolaris species specific region of 28S gene.

- (iii) HYPOTHETICAL: No
- (iv) ANTISENSE: No
 - (v) SEQUENCE DESCRIPTION: SEQ ID NO: 36:

AGCCAGACTT GCTTGCAGTT GCTCATCCGG GCTTTTGCCC
GGTGCACTCT 50

TCTGCAGGCA GGCCAGCATC AGTTTGGGCG GTGGGATAAA GGTCTCTGTC 100

ACGTACCTTC CTTCGGGTTG GCCATATAGG GGAGACGTCA
TACCACCAGC 150

CTGGACTGAG GTCCGCGCAT CTGCTAGGAT GCTGGCGTAA
TGGCTGTAAG 200

CGGCCCGTCT TGA

213

(38) INFORMATION FOR SEQ ID NO 37:

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- (i) SEQUENCE CHARACTERISTICS:
 - (A) LENGTH: 105
 - (B) TYPE: nucleic acid
 - (C) STRANDEDNESS: single
 - (D) TOPOLOGY: linear

(ii) MOLECULE TYPE: Blastoschizomyces species specific region of 28S gene.

- (iii) HYPOTHETICAL: No
- (iv) ANTISENSE: No
- (v) SEQUENCE DESCRIPTION: SEQ ID NO: 37:

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TGAAATTGTT GAAAGGGAAG GCGATGGTAG GAATAAGAGG CTGCGGTTTG 50

AAATAATTGT TTTTCGGGCC ACGGTCTCCT GAGCCTGCTT
TCGCACÇCGT 100

CTTGA

105

- (39) INFORMATION FOR SEQ ID NO 38:
 - (i) SEQUENCE CHARACTERISTICS:
 - (A) LENGTH: 214
 - (B) TYPE: nucleic acid
 - (C) STRANDEDNESS: single
 - (D) TOPOLOGY: linear

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- (ii) MOLECULE TYPE: Blastomyces dermatitidis specific region of 28S gene.
- (iii) HYPOTHETICAL: No

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- (iv) ANTISENSE: No
- (v) SEQUENCE DESCRIPTION: SEQ ID NO: 38:

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GACCAGAGTC GGCCGTGGGG GTTCAGCGGG CATTCGTTGC CCGTGCACTC 50

CCCCACGGC GGGCCAGCGT CGGTTTCGAC GGCCGGTCAA
AGGCCCCCGG 100

AATGTGTCGC CTCTCGGGGC GTCTTATAGC CGGGGGTGCA
ATGCGGCCAG 150

WO 98/55649 PCT/IB98/00865

TCGGGACCGA GGAACGCGCT TCGGCACGGA CGCTGGCTTA ATGGTCGTAA 200

GCGACCCGTC TTGA

214

(40) INFORMATION FOR SEO ID NO 39:

- (i) SEQUENCE CHARACTERISTICS:
 - (A) LENGTH: 213
 - (B) TYPE: nucleic acid.
 - (C) STRANDEDNESS: single
 - (D) TOPOLOGY: linear
- (ii) MOLECULE TYPE: Chrysosporium species specific region of 28S gene.
 - (iii) HYPOTHETICAL: No
 - (iv) ANTISENSE: No

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(v) SEQUENCE DESCRIPTION: SEQ ID NO: 39:

AACCAGACTT GCGCGCGGCC GATCATCCGG TGTTCTCACC
GGTGCACTCG 50

GCCGTGCTCA GGCCAGCATC GGTTTTGGCG GCTGGATAAA GGCCCTAGGA 100

ATGTGGCTCC TCTCGGGGAG TGTTATAGCC TAGGGTGCAA
TGCAGCCTGC 150

TGGGACCGAG GACCGCGCTT CGGCTAGGAT GCTGGCGTAA TGGTTGTAAG 200

CGGCCCGTCT TGA

(41) INFORMATION FOR SEQ ID NO 40:

- (i) SEQUENCE CHARACTERISTICS:
 - (A) LENGTH: 207
 - (B) TYPE: nucleic acid
 - (C) STRANDEDNESS: single
 - (D) TOPOLOGY: linear

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- (ii) MOLECULE TYPE: Cladosporium species specific region of 28S gene.
- (iii) HYPOTHETICAL: No
- (iv) ANTISENSE: No
- (v) SEQUENCE DESCRIPTION: SEQ ID NO: 40:
- 20 AACCAGACTT GCTCGCGGTG TTCCGCCGGT CTTCTGACCG
 GTCTACTCGC 50

CGCGTTGCAG GCCAGCATCG TCTGGTGCCG CTGGATAAGA
CTTGAGGAAT 100

GTAGCTCCCT CGGGAGTGTT ATAGCCTCTT GTGATGCAGC
GAGCGCCGGG 150

CGAGGTCCGC GCTTCGGCTA GGATGCTGGC GTAATGGTCG
TAATCCGCCC 200

GTCTTGA

207

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(42) INFORMATION FOR SEQ ID NO 41:

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- (i) SEQUENCE CHARACTERISTICS:
 - (A) LENGTH: 213
 - (B) TYPE: nucleic acid
 - (C) STRANDEDNESS: single
 - (D) TOPOLOGY: linear
- (ii) MOLECULE TYPE: Curvularia species specific region of 28S gene.
- (iii) HYPOTHETICAL: No

(iv) ANTISENSE: No

(v) SEQUENCE DESCRIPTION: SEQ ID NO: 41:

AGCCAGACTT GCTTGCAGTT GCTCATCCGG GCTTTTGCCC
GGTGCACTCT 50

TCTGCAGGCA GGCCAGCATC AGTTTGGGCG GTGGGATAAA GGTCTCTGAC 100

ACGTTCCTTC CTTCGGGTTG GCCATATAGG GGAGACGTCA
TACCACCAGC 150

CTGGACTGAG GTCCGCGCAT CTGCTAGGAT GCTGGCGTAA
TGGCTGTAAG 200

CGGCCCGTCT TGA

213

(43) INFORMATION FOR SEQ ID NO 42:

- (i) SEQUENCE CHARACTERISTICS:
 - (A) LENGTH: 213
 - (B) TYPE: nucleic acid
 - (C) STRANDEDNESS: single

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(D) TOPOLOGY: linear

- (ii) MOLECULE TYPE: Candida albicans specific region of 28S gene.
- (iii) HYPOTHETICAL: No
- (iv) ANTISENSE: No
- (v) SEQUENCE DESCRIPTION: SEQ ID NO: 42:

GATCAGACTT GGTATTTTGC ATGCTGCTCT CTCGGGGGCG GCCGCTGCGG 50

TTTACCGGGC CAGCATCGGT TTGGAGCGGC AGGATAATGG
CGGAGGAATG 100

TGGCACGCT TCTGCTGTGT GTTATAGCCT CTGACGATGC
TGCCAGCCTA 150

GACCGAGGAC TGCGGTTTTT AACCTAGGAT GTTGGCATAA
TGATCTTAAG 200

20 TCGCCCGTCT TGA

- (44) INFORMATION FOR SEQ ID NO 43:
- (i) SEQUENCE CHARACTERISTICS:
 - (A) LENGTH: 223
 - (B) TYPE: nucleic acid
 - (C) STRANDEDNESS: single
 - (D) TOPOLOGY: linear
 - (ii) MOLECULE TYPE: Candida glabrata specific region of 28S gene.

(III) HYP(OTHETIC	:AL: No	
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- (iv) ANTISENSE: No
- (v) SEQUENCE DESCRIPTION: SEQ ID NO: 43:

GATCAGACAT GGTGTTTTGC GCCCCTTGCC TCTCGTGGGC
TTGGGACTCT 50

CGCAGCTCAC TGGGCCAGCA TCGGTTTTGG CGGCCGGAAA
AAACCTAGGG 100

AATGTGGCTC TGCGCCTCGG TGTAGAGTGT TATAGCCCTG
GGGAATACGG 150

CCAGCCGGGA CCGAGGACTG CGATACTTGT TATCTAGGAT
GCTGGCATAA 200

TGGTTATATG CCGCCCGTCT TGA

223

(45) INFORMATION FOR SEQ ID NO 44:

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- (i) SEQUENCE CHARACTERISTICS:
 - (A) LENGTH: 212
 - (B) TYPE: nucleic acid
 - (C) STRANDEDNESS: single
 - (D) TOPOLOGY: linear
- (ii) MOLECULE TYPE: Candida guilliermondii specific region of 28S gene.
- (iii) HYPOTHETICAL: No

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(iv) ANTISENSE: No

(v) SEQUENCE DESCRIPTION: SEQ ID NO: 44:

GATCAGACTC GATATTTTGT GAGCCTTGCC TTCGTGGCGG GGTGACCCGC 50

AGCTTATCGG GCCAGCATCG GTTTGGGCGG TAGGATAATG GCGTAGGAAT 100

GTGACTTTRC TTCGGTGAAG TGTTATAGCC TGCGTTGATG
CTGCCTGCCT 150

AGACCGAGGA CTGCGATTTT ATCAAGGATG CTGGCATAAT
GATCCCAAAC 200

CGCCCGTCTT GA

212

(46) INFORMATION FOR SEQ ID NO 45:

- (i) SEQUENCE CHARACTERISTICS:
 - (A) LENGTH: 214
 - (B) TYPE: nucleic acid
 - (C) STRANDEDNESS: single
 - (D) TOPOLOGY: linear
- (ii) MOLECULE TYPE: Coccidioides immitis specific region of 28S gene.
- 25 (iii) HYPOTHETICAL: No
 - (iv) ANTISENSE: No
 - (v) SEQUENCE DESCRIPTION: SEQ ID NO: 45:

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AACCAGACTC GGTCGTGGGG GCTCAGCGGG CATGAGTGCC CGTGTACTCC 50

CCCATGCTCC GGGCCAGCAT CAGTTCTGGC GGTTGGTTAA
AGGCCTCTGG 100

AATGTATCGT CCTCCGGGAC GTCTTATAGC CAGGGGCGCA
ATGCGGCCAG 150

CCGGGACTGA GGAACGCGCT TCGGCACGGA TGCTGGCATA ATGGTTGTAA 200

GCGGCCCGTC TTGA

214

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(47) INFORMATION FOR SEQ ID NO 46:

- (i) SEQUENCE CHARACTERISTICS:
 - (A) LENGTH: 187
 - (B) TYPE: nucleic acid
 - (C) STRANDEDNESS: single
 - (D) TOPOLOGY: linear
- (ii) MOLECULE TYPE: Candida kefyr specific region of 28S gene.
- (iii) HYPOTHETICAL: No
- (iv) ANTISENSE: No

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(v) SEQUENCE DESCRIPTION: SEQ ID NO: 46:

GATCAGACAT GGCGTTTGCT TCGGCTTTCG CTGGGCCAGC

30 ATCAGTTTA 50

GCGGTTGGAT AAATCCTCGG GAATGTGGCT CTGCTTCGGT
AGAGTGTTAT 100

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AGCCCGTGGG AATACAGCCA GCTGGGACTG AGGATTGCGA CTTTTGTCAA 150

GGATGCTGGC GTAATGGTTA AATGCCGCCC GTCTTGA

187

(48) INFORMATION FOR SEQ ID NO 47:

- (i) SEQUENCE CHARACTERISTICS:
 - (A) LENGTH: 213
 - (B) TYPE: nucleic acid
 - (C) STRANDEDNESS: single
 - (D) TOPOLOGY: linear
- (ii) MOLECULE TYPE: Candida krusei specific region of 28S gene.
- (iii) HYPOTHETICAL: No
- (iv) ANTISENSE: No
- (v) SEQUENCE DESCRIPTION: SEQ ID NO: 47:

CGCCCGACAT GGGGATTGCG CACCGCTGCC TCTCGTGGGC
GGCGCTCTGG 50

GCTTTCCCTG GGCCAGCATC GGTTCTTGCT GCAGGAGAAG GGGTTCTGGA 100

ACGTGGCTCT TCGGAGTGTT ATAGCCAGGG CCAGATGCTG CGTGCGGGA 150

CCGAGGACTG CGGCCGTGTA GGTCACGGAT GCTGGCAGAA

CGGCGCAACA 200

CCGCCCGTCT TGA

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(49) INFORMATION FOR SEQ ID NO 48:

- (i) SEQUENCE CHARACTERISTICS:
 - (A) LENGTH: 236
 - (B) TYPE: nucleic acid
 - (C) STRANDEDNESS: single
 - (D) TOPOLOGY: linear
- (ii) MOLECULE TYPE: Cryptococcus laurentii specific region of 28S gene.
 - (iii) HYPOTHETICAL: No
 - (iv) ANTISENSE: No
 - (v) SEQUENCE DESCRIPTION: SEQ ID NO: 48:

AGTCAGTCGT GTCTGGGAGG CTCAGCCGGT TCTGCCGGTG

20 TATTCCTCTC 50

AGACGGGTCA ACATCAGTTT TGTCCGACGG ATAATGGCGG CGGGAAAGTA 100

GCACCTCCGG GTGTGTTATA GCCCGCTGTC GCATACGCCG
GATGAGACTG 150

25 AGGCATGCAG CTCGCCTTTA TGGCAGGGGT TCGCCCACTT
TCGAGCTTAG 200

GATGTTGACG TAATGGCTTT AAACGACCCG TCTTGA

- (50) INFORMATION FOR SEQ ID NO 49:
 - (i) SEQUENCE CHARACTERISTICS:

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(A) LENGTH: 173

(B) TYPE: nucleic acid

(C) STRANDEDNESS: single

(D) TOPOLOGY: linear

(ii) MOLECULE TYPE: Candida lusitaniae specific region of 28S gene.

(iii) HYPOTHETICAL: No

(iv) ANTISENSE: No

(v) SEQUENCE DESCRIPTION: SEQ ID NO: 49:

AAGCAGACAC GGTTTTACCG GGCCAGCGTC GAAAAGGGGG GAGGAACAAG 50

AACTCGAGAA TGTGGCGCGC ACCTTCGGGY GCGCGTGTTA
TAGCTCGTGT 100

TGACGCCTCC ATCCCTTTTC GAGGCCTGCG ATTCTAGGAC
GCTGGCGTAA 150

TGGTTGCAAG CCGCCGTCT TGA

- (51) INFORMATION FOR SEQ ID NO 50:
 - (i) SEQUENCE CHARACTERISTICS:
 - (A) LENGTH: 238
 - (B) TYPE: nucleic acid
 - (C) STRANDEDNESS: single
 - (D) TOPOLOGY: linear
 - (ii) MOLECULE TYPE: Cryptococcus neoformans var gattii (serotype B)

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specific region of 28S gene.

- (iii) HYPOTHETICAL: No
- (iv) ANTISENSE: No
- (v) SEQUENCE DESCRIPTION: SEQ ID NO: 50:

AGTCAGTCGT GTCTATTGGG TTCAGCCAGC TCTGCTGGTG

TATTCCCTTT 50

AGACGGGTCA ACATCAGTTC TGATCGGTGG ATAAGGGCTG
GAGGAATGTG 100

GCACTCTTCG GGGTGTGTTA TAGCCTCCTG TCGCATACAC
TGGTTGGGAC 150

TGAGGAATGC AGCTCGCCTT TATGGCCGGG GTTCGCCCAC
GTTCGAGCTT 200

AGGATGTTGA CAAAATGGCT TTAAACGACC CGTCTTGA 238

- (52) INFORMATION FOR SEQ ID NO 51:
 - (i) SEQUENCE CHARACTERISTICS:
 - (A) LENGTH: 238
 - (B) TYPE: nucleic acid
 - (C) STRANDEDNESS: single
 - (D) TOPOLOGY: linear
 - (ii) MOLECULE TYPE: Cryptococcus neoformans (serotype A) specific region
- of 28S gene.
 - (iii) HYPOTHETICAL: No

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- (iv) ANTISENSE: No
- (v) SEQUENCE DESCRIPTION: SEQ ID NO: 51:

AGTCAGTCGT GTCTATTGGG TTCAGCCAGT TCTGCTGGTG
TATTCCCTTT 50

AGACGGGTCA ACATCAGTTC TGATCGGTGG ATAAGGGCTG GGGGAATGTA 100

GCACTCTTCG GAGTGTGTTA TAGCCTCCTG TCGCATACAC
TGGTTGGGAC 150

TGAGGAATGC AGCTCGCCTT TATGGCCGGG GTTCGCCCAC GTTCGAGCTT 200

AGGATGTTGA CAAAATGGCT TTAAACGACC CGTCTTGA
238

- (53) INFORMATION FOR SEQ ID NO 52:
 - (i) SEQUENCE CHARACTERISTICS:
 - (A) LENGTH: 211
 - (B) TYPE: nucleic acid
 - (C) STRANDEDNESS: single
 - (D) TOPOLOGY: linear
 - (ii) MOLECULE TYPE: Candida parapsilosis specific region of 28S gene.
 - (iii) HYPOTHETICAL: No
 - (iv) ANTISENSE: No

(v) SEQUENCE DESCRIPTION: SEQ ID NO: 52:

GATCAGACTT GGTATTTTGT ATGTTACTCT CTCGGGGGTG
GCCTCTACAG 50

TTTACCGGGC CAGCATCAGT TTGAGCGGTA GGATAAGTGC AAAGAAATGT 100

GGCACTGCTT CGGTAGTGTG TTATAGTCTT TGTCGATACT
GCCAGCTTAG 150

ACTGAGGACT GCGGCTTCGG CCTAGGATGT TGGCATAATG
ATCTTAAGTC 200

GCCCGTCTTG A

211

(54) INFORMATION FOR SEQ ID NO 53:

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- (i) SEQUENCE CHARACTERISTICS:
 - (A) LENGTH: 238
 - (B) TYPE: nucleic acid
 - (C) STRANDEDNESS: single
 - (D) TOPOLOGY: linear
- (ii) MOLECULE TYPE: Cryptococcus terreus specific region of 28S gene.
- (iii) HYPOTHETICAL: No

- (iv) ANTISENSE: No
- (v) SEQUENCE DESCRIPTION: SEQ ID NO: 53:
- AGTCAGTCAT GTCTATTGGA CTCAGCCGGT TCTGCCGGTG

 TACTTCCTTT 50

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AGATGGGGTC AACATCAGTT TTGATCGCTG GAAAAGGGCA GGAGGAATGT 100

AGCACTCTCG GGTGAACTTA TAGCCTTCTG TCGTATACAG
TGGTTGGGAC 150

TGAGGAACGC AGCATGCCTT TATGGCCGGG GTTCGCCCAC
GTACATGCTT 200

AGGATGTTGA CATAATGGCT TTAAACGACC CGTCTTGA
238

- (55) INFORMATION FOR SEQ ID NO 54:
 - (i) SEQUENCE CHARACTERISTICS:
 - (A) LENGTH: 211
 - (B) TYPE: nucleic acid
 - (C) STRANDEDNESS: single
 - (D) TOPOLOGY: linear
 - (ii) MOLECULE TYPE: Candida tropicalis specific region of 28S gene.
- 20 (iii) HYPOTHETICAL: No

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- (iv) ANTISENSE: No
- (v) SEQUENCE DESCRIPTION: SEQ ID NO: 54:

GATCAGACTT GGTATTTTGT ATGTTACTTC TTCGGGGGTG
GCCTCTACAG 50

TTTATCGGGC CAGCATCAGT TTGGGCGGTA GGAGAATTGC GTTGGAATGT 100

GGCACGGCTT CGGTTGTGTG TTATAGCCTT CGTCGATACT
GCCAGCCTAG 150

ACTGAGGACT GCGGTTTATA CCTAGGATGT TGGCATAATG ATCTTAAGTC 200

GCCCGTCTTG A

211

- (56) INFORMATION FOR SEQ ID NO 55:
 - (i) SEQUENCE CHARACTERISTICS:
 - (A) LENGTH: 211
 - (B) TYPE: nucleic acid
 - (C) STRANDEDNESS: single
 - (D) TOPOLOGY: linear
 - (ii) MOLECULE TYPE: Fusarium species specific region of 28S gene.
 - (iii) HYPOTHETICAL: No
 - (iv) ANTISENSE: No
 - (v) SEQUENCE DESCRIPTION: SEQ ID NO: 55:

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GACCAGACTT GGGCTTGGTT AATCATCTGG GGTTCTCYCC AGTGCACTTT 50

TCCAGTCCAG GCCAGCATCA GTTTTCSCCG GGGGATAAAG
RCTTCGGGAA 100

TGTGGCTCYC YYCGGGGAGT GTTATAGCCC GTTGYGTAAT ACCCTGGBGG 150

GGACTGAGGT TCGCGCWTCT GCAAGGATGC TGGCGTAATG GTCATCAACG 200

o ACCCGTCTTG A

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(57) INFORMATION FOR SEQ ID NO 56:

- (i) SEQUENCE CHARACTERISTICS:
 - (A) LENGTH: 238
 - (B) TYPE: nucleic acid
 - (C) STRANDEDNESS: single
 - (D) TOPOLOGY: linear
- (ii) MOLECULE TYPE: Filobasidium capsuligenum specific region of 28S gene.
 - (iii) HYPOTHETICAL: No
 - (iv) ANTISENSE: No
 - (v) SEQUENCE DESCRIPTION: SEQ ID NO: 56:

AGTCAGTCAT GTCTATTGGA CTCAGCCGGT TCTGCCGGTG

20 TATTTCCTTT 50

AGATGGGGTC AACATCAGTT TTGACCGTTG GATAAAGGCA GGAAGAATGT 100

AGCACTCTCG GGTGAACTTA TAGCTTCTTG TCACATACAA
TGGTTGGGAC 150

TGAGGAACGC AGCATGCCTT TATGGCCGGG ATTCGTCCAC
GTACATGCTT 200

AGGATGTTGA CATAATGGCT TTAAACGACC CGTCTTGA
238

(58) INFORMATION FOR SEQ ID NO 57:

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- (i) SEQUENCE CHARACTERISTICS:
 - (A) LENGTH: 238
 - (B) TYPE: nucleic acid
 - (C) STRANDEDNESS: single
 - (D) TOPOLOGY: linear
- (ii) MOLECULE TYPE: Filobasidiella neoformans var bacillispora (serotype
- C) specific region of 28S gene.
- (iii) HYPOTHETICAL: No
- (iv) ANTISENSE: No
- (v) SEQUENCE DESCRIPTION: SEQ ID NO: 57:

AGTCAGTCGT GTCTATTGGG TTCAGCCAGC TCTGCTGGTG TATTCCCTTT 50

AGACGGGTCA ACATCAGTTC TGATCGGTGG ATAAGGGCTG
GAGGAATGTG 100

GCACTCTTCG GGGTGTGTTA TAGCCTCCTG TCGCATACAC
TGGTTGGGAC 150

TGAGGAATGC AGCTCGCCTT TATGGCCGGG GTTCGCCCAC GTTCGAGCTT 200

AGGATGTTGA CAAAATGGCT TTAAACGACC CGTCTTGA
238

- (59) INFORMATION FOR SEQ ID NO 58:
 - (i) SEQUENCE CHARACTERISTICS: (A) LENGTH: 238

- (B) TYPE: nucleic acid
- (C) STRANDEDNESS: single
- (D) TOPOLOGY: linear
- (ii) MOLECULE TYPE: Filobasidiella neoformans var neoformans (serotype D) specific region of 28S gene.
- (iii) HYPOTHETICAL: No
- (iv) ANTISENSE: No
 - (v) SEQUENCE DESCRIPTION: SEQ ID NO: 58:
- AGTCAGTCGT GTCTATTGGG TTCAGCCAGT TCTGCTGGTG
 TATTCCCTTT 50

AGACGGGTCA ACATCAGTTC TGATCGGTGG ATAAGGGCTG
GAGGAATGTG 100

GCACTCTTCG GGGTGTGTTA TAGCCTCCTG TCGCATACAC
TGGTTGGGAC 150

TGAGGAATGC AGCTCGCCTT TATGGCCGGG GTTCGCCCAC
GTTCGAGCTT 200

AGGATGTTGA CAAAATGGCT TTAAACGACC CGTCTTGA
238

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- (60) INFORMATION FOR SEQ ID NO 59:
 - (i) SEQUENCE CHARACTERISTICS:
 - (A) LENGTH: 236
 - (B) TYPE: nucleic acid
 - (C) STRANDEDNESS: single

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(D) TOPOLOGY: linear

- (ii) MOLECULE TYPE: Filobasidium uniguttulatum specific region of 28S gene.
- (iii) HYPOTHETICAL: No
- (iv) ANTISENSE: No
- (v) SEQUENCE DESCRIPTION: SEQ ID NO: 59:

AGTCAGTCGT GCTCAATGGA CTCAGCCGTT CTGCGGTGTA
TTTCCATTGG 50

GTGGGGTCAA CATCAGTTTT GATCGCTGGA TAAAGGCAGG AGGAATGTAG 100

CACCCCGGG TGAACTTATA GCCTCTTGTC ACATACAGTG
GTTGGGACTG 150

AGGAACGCAG CATGCCTTTA TGGCCGGGAT TCGTCCACGT

20 ACATGCTTAG 200

GATGTTGACA TAATGGCTTT AAACGACCCG TCTTGA

- (61) INFORMATION FOR SEQ ID NO 60:
 - (i) SEQUENCE CHARACTERISTICS:
 - (A) LENGTH: 204
 - (B) TYPE: nucleic acid
 - (C) STRANDEDNESS: single
- (D) TOPOLOGY: linear.
 - (ii) MOLECULE TYPE: Geotrichum species specific region of 28S gene.

- (iii) HYPOTHETICAL: No
- (iv) ANTISENSE: No
- (v) SEQUENCE DESCRIPTION: SEQ ID NO: 60:

AATCAGACTT GGTGCTGTTG TTCAACTRTG TTTCGGCATA

GTGTACTCAG 50

CAGTACTAGG CCAAGGTGGG GTGTTTGGGA GTGAAAAAGA AGTAGGAACG 100

TAACTCTTCG GAGTGTTATA GCCTACTTTC ATAGCTCCTC AGGCGCCTCA 150

GGACTGCGCT TCGGCAAGGA CCTTGGCATA ATGATTCTAT
ACCGCCCGTC 200

TTGA

204

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- (62) INFORMATION FOR SEQ ID NO 61:
 - (i) SEQUENCE CHARACTERISTICS:
 - (A) LENGTH: 214
 - (B) TYPE: nucleic acid
 - (C) STRANDEDNESS: single
 - (D) TOPOLOGY: linear
 - (ii) MOLECULE TYPE: Histoplasma capsulatum specific region of 28S gene.
 - (iii) HYPOTHETICAL: No

(iv) ANTISENSE: No

(v) SEQUENCE DESCRIPTION: SEQ ID NO: 61:

GAYCAGAGTC GGCCGYGGGG GTTCAGCGGG CATTCGTTGC CCGTGCAATC 50

CCCCGCGGCC GGGCCAGCGT CGGTTTCGAC GGCCGGTCAA
AGGCCCCCGG 100

AATGTGTCGC CTCTCGGGGC GTCTTATAGC CGGGGGTGCA
ATGCGGCCAG 150

TCGGGACCGA GGAACGCGCT CCGGCACGGA CGCTGGCTTA
ATGGTCGTCA 200

GCGACCCGTC TTGA

214 -

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(63) INFORMATION FOR SEQ ID NO 62:

- (i) SEQUENCE CHARACTERISTICS:
 - (A) LENGTH: 215
 - (B) TYPE: nucleic acid
 - (C) STRANDEDNESS: single
 - (D) TOPOLOGY: linear
- (ii) MOLECULE TYPE: Malbranchea species specific region of 28S gene.
- (iii) HYPOTHETICAL: No
- (iv) ANTISENSE: No

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(v) SEQUENCE DESCRIPTION: SEQ ID NO: 62:

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AGACAGACTC GAGCGCGGG GCTCAGCGGG TATTGTTATG
CCCGTGCACT 50

CCCCGCGCC CGGGCCAGCA TCAGTTTTGG CGGCCGGTCA
AAGGCCCTTG 100

GAATGTATCG TCCTCCGGGA CGTCTTATAG CCAAGGGTGC
AATGCGGCCA 150

GCCGGGACTG AGGAACGCGC TTCGGCACGG ATGCTGGCGT
AATGGCTGTA 200

AGCGGCCCGT CTTGA

215

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(64) INFORMATION FOR SEQ ID NO 63:

- (i) SEQUENCE CHARACTERISTICS:
 - (A) LENGTH: 237
 - (B) TYPE: nucleic acid
 - (C) STRANDEDNESS: single
 - (D) TOPOLOGY: linear
- (ii) MOLECULE TYPE: Mucor species specific region of 28S gene.
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- (iii) HYPOTHETICAL: No
- (iv) ANTISENSE: No
- (v) SEQUENCE DESCRIPTION: SEQ ID NO: 63:

AGCCAGACTG GTTTGACTGT AATCAACCTA GAATTCGTTC
TGGGTGCACT 50

TGCAGTCTAT ACCTGCCAAC AACAGTTTGA TTTGGAGGAA
AAAATTAGTA 100

GGAATGTAGC CTCTCGAGGT GTTATAGCCT ACTATCATAC
TCTGGATTGG 150

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ACTGAGGAAC GCAGCGAATG CCWTTAGGCR AGATTGCTGG GTGCTTTCGC 200

TAATAAATGT TAGAATTTCT GCTTCGGGTG GTGCTAA

237

(65) INFORMATION FOR SEQ ID NO 64:

- (i) SEQUENCE CHARACTERISTICS:
 - (A) LENGTH: 209
 - (B) TYPE: nucleic acid
 - (C) STRANDEDNESS: single
 - (D) TOPOLOGY: linear
- (ii) MOLECULE TYPE: Paecilomyces species specific region of 28S gene.
- (iii) HYPOTHETICAL: No
- (iv) ANTISENSE: No

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(v) SEQUENCE DESCRIPTION: SEQ ID NO: 64:

GACCAGACTT GGGCCCGGTG GATCATCCAG CGTTCTCGCT
GGTGCACTCC 50

GCCGGGTTCA GGCCAGCATC AGTTCGCCGC GGGGGAAAAA
GGCTTCGGGA 100

ACGTGGCTCC TACGGGAGTG TTATAGCCCG TTGCATAATA
CCCTGGGGCG 150

GACTGAGGTT CGCGCTCCGC AAGGATGCTG GCGTAATGGT CATCAGCGAC 200

30 CCGTCTTGA 209

(66) INFORMATION FOR SEQ ID NO 65:

- (i) SEQUENCE CHARACTERISTICS:
 - (A) LENGTH: 199
 - (B) TYPE: nucleic acid
 - (C) STRANDEDNESS: single
 - (D) TOPOLOGY: linear

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- (ii) MOLECULE TYPE: Penicillium species specific region of 28S gene.
- (iii) HYPOTHETICAL: No
- (iv) ANTISENSE: No
- (v) SEQUENCE DESCRIPTION: SEQ ID NO: 65:
- 20 GACCAGACTC GCCCACGGGG TTCAGCCGGC ATTCGTGCCG
 GTGTACTTCC 50

CCGCGGCGG GCCAGCGTCG GTTTGGKCGG CCGGTCAAAG GCCCTCGGAA 100

TRTAACGCCC CCCGGGGCGT CTTATAGCCG AGGGTGCCAT
25 GCGGCCAGCM 150

CAGACCGAGG AACGCGCTTC GGCTCGGACG CTGGCATAAT GGTCGTAAA 199

- 30 (67) INFORMATION FOR SEQ ID NO 66:
 - (i) SEQUENCE CHARACTERISTICS:

(A) LENGTH: 210

(B) TYPE: nucleic acid

(C) STRANDEDNESS: single

(D) TOPOLOGY: linear

(ii) MOLECULE TYPE: Pseudallescheria boydii region of 28S gene.

(iii) HYPOTHETICAL: No

(iv) ANTISENSE: No

(v) SEQUENCE DESCRIPTION: SEQ ID NO: 66:

GACCAGACTT GTGCCCGTCG AATCAGCCGC CGCTCGTCGG
CGGCGCACTT 50

CGGCGGCTC AGGCCAGCAT CAGTTCGCTG CAGGGGGAGA
AAGGCGATGG 100

GAATGTGGCT CTTCGGAGTG TTATAGCCCG CCGCGCAATA
CCCCTCGGCG 150

GACTGAGGAC CGCGCATCTG CAAGGATGCT GGCGTAATGG TCGTCAGCGA 200

CCCGTCTTGA 210

(68) INFORMATION FOR SEQ ID NO 67:

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(i) SEQUENCE CHARACTERISTICS:

(A) LENGTH: 244

(B) TYPE: nucleic acid

(C) STRANDEDNESS: single

(D) TOPOLOGY: linear

- (ii) MOLECULE TYPE: Rhizopus species (NO: 1) specific region of 28S gene.
- (iii) HYPOTHETICAL: No
- (iv) ANTISENSE: No
- (v) SEQUENCE DESCRIPTION: SEQ ID NO: 67:

AGCCAGACTG GCTTGTCTGT AATCAATCTA GGTTTCGTGC CTGGATGCAC 50

TTGCAGACTA TTTGCCTGCC AACGACAATT TTTTTTGAGT
GTAAAAACTA 100

TTGGAAATGT GGCCAATATT TATTTATTGG TGTTATAGTC
CTTTAGAAAA 150

TACCTTGAAT TGGATTGAGG AACGCAGCGA ATGCTTCTCT
TTNGAGGCAA 200

AGTCTTTAT TGGGATTTAC GGATCAGACT GTGGCATTGT CACA

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- (69) INFORMATION FOR SEQ ID NO 68:
 - (i) SEQUENCE CHARACTERISTICS:
 - (A) LENGTH: 215
 - (B) TYPE: nucleic acid
 - (C) STRANDEDNESS: single
 - (D) TOPOLOGY: linear
- (ii) MOLECULE TYPE: Rhizopus species (NO: 2) specific region of 28S gene.

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- (iii) HYPOTHETICAL: No
- (iv) ANTISENSE: No
- (v) SEQUENCE DESCRIPTION: SEQ ID NO: 68:

AGCCAGACTG GCTTGTCTGT AATCAATCTA GGCTTCGGCC
TGGATGCACT 50

TGCAGGCTAT GCCTGCCAAC GACAATTTGA CTTGAGGGAA

AAAACTAGGG 100

GAAATGTGGC CCACTTGTGG GTGTTATAGT CCCTTAGAAA ATACCTTGGG 150

TTGGATTGAG GAACGCAGCG AATGCTTATT GGCGAGTTTT
CCAGGAAGGT 200

TTTCTGAGGT ACTAC

215

(70) INFORMATION FOR SEQ ID NO 69:

- (i) SEQUENCE CHARACTERISTICS:
 - (A) LENGTH: 215
 - (B) TYPE: nucleic acid
 - (C) STRANDEDNESS: single
 - (D) TOPOLOGY: linear
- (ii) MOLECULE TYPE: Rhizopus species (NO: 3) specific region of 28S gene.
 - (iii) HYPOTHETICAL: No
- 30 (iv) ANTISENSE: No
 - (v) SEQUENCE DESCRIPTION: SEQ ID NO: 69:

AGCCAGACTG GCTTGTCTGT AATCAGTCTA AGCTTCGGCT TGGATGCACT 50

TGCAGGCTAT GCCTGCCAAC GACAATTTGG CTTGAGGGAA
AAAACTAAGG 100

GAAATGTGGC CCATCCGTGG GTGTTATAGT CCCTTAGAAA ATACCTTGGG 150

CTGGATTGAG GTACGCAGCG AATGCTATTT GGCGAGTTGG CTGGGAATAT 200

TTTCTGAGGT GCTTT

215

(71) INFORMATION FOR SEQ ID NO 70:

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- (i) SEQUENCE CHARACTERISTICS:
 - (A) LENGTH: 210
 - (B) TYPE: nucleic acid
 - (C) STRANDEDNESS: single
 - (D) TOPOLOGY: linear
- (ii) MOLECULE TYPE: Sporothrix species specific region of 28S gene.
- (iii) HYPOTHETICAL: No

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- (iv) ANTISENSE: No
- (v) SEQUENCE DESCRIPTION: SEQ ID NO: 70:

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GACCAGACTT GCGCCYCGCG GACCACCCGG CGTTCTCGCC
GGTGCACTCT 50

GCGKKGCGCA GGCCAGCATC GGTTCTCCCA GGGGGACAAA GGCCGCGGA 100

ACGTAGCTCC TTCGGGAGTG TTATAGCCCG CGGCGGCATG CCCCTGGGGG 150

GACCGAGGAC CGCGCTTCGG CAAGGATGCT GGCGTAATGG TCACCAGCGA 200

ACCGTCTTGA

210

- (72) INFORMATION FOR SEQ ID NO 71:
 - (i) SEQUENCE CHARACTERISTICS:
 - (A) LENGTH: 208
 - (B) TYPE: nucleic acid
 - (C) STRANDEDNESS: single
 - (D) TOPOLOGY: linear
 - (ii) MOLECULE TYPE: Scopulariopsis brevicaulis specific region of 28S gene.

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- (iii) HYPOTHETICAL: No
- (iv) ANTISENSE: No
- (v) SEQUENCE DESCRIPTION: SEQ ID NO: 71:

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GACCAGACTT GCGCCCGTCG GATCAACCGT CGCTTGCGGC
GGCGCACTCC 50

GGCGGCTCA GGCCAGCATC AGTTCGTCCG GGGGGAGAAA GGCGGCGGA 100

ATGTGGCTCT TCGGAGTGTT ATAGCCCGCC GTGTAATACC
CTCGGGTGGA 150

CTGAGGACCG CGCGTATGCA AGGATGCTGG CGTAATGGTC GTCAGCGACC 200

CGTCTTGA

208

- (73) INFORMATION FOR SEQ ID NO 72:
 - (i) SEQUENCE CHARACTERISTICS:
 - (A) LENGTH: 210
 - (B) TYPE: nucleic acid
 - (C) STRANDEDNESS: single
 - (D) TOPOLOGY: linear
 - (ii) MOLECULE TYPE: Scopulariopsis brumptii specific region of 28S gene.
 - (iii) HYPOTHETICAL: No
 - (iv) ANTISENSE: No
 - (v) SEQUENCE DESCRIPTION: SEQ ID NO: 72:

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GACCAGACTC GCGCCCGTCG GATCAGCCGT CGCTCGTCGG
CGGCGCACTC 50

CGGCGGCTC GGGCCAGCAT CAGTTCGCCT CGGGGGGAGA
AAGGCGGCGG 100

GAATGTGGCT CTACGGAGTG TTATAGCCCG CCGCGTAATA
CCCCCGGGCG 150

GACTGAGGAC CGCGCGTATG CAAGGATGCT GGCGTAATGG
TCGTCAGCGA 200

o CCCGTCTTGA

(74) INFORMATION FOR SEQ ID NO 73:

- (i) SEQUENCE CHARACTERISTICS:
 - (A) LENGTH: 214
 - (B) TYPE: nucleic acid
 - (C) STRANDEDNESS: single
 - (D) TOPOLOGY: linear
- (ii) MOLECULE TYPE: Saccharomyces cerevisiae specific region of 28S gene.
 - (iii) HYPOTHETICAL: No
 - (iv) ANTISENSE: No
 - (v) SEQUENCE DESCRIPTION: SEQ ID NO: 73:
- 20 GATCAGACAT GGTGTTTTGT GCCCTCTGCT CCTTGTGGGT
 AGGGGAATCT 50

CGCATTTCAC TGGGCCAGCA TCAGTTTTGG TGGCAGGATA
AATCCATAGG 100

AATGTAGCTT GCCTCGGTAA GTATTATAGC CTGTGGGAAT
ACTGCCAGCT 150

GGGACTGAGG ACTGCGACGT AAGTCAAGGA TGCTGGCATA
ATGGTTATAT 200

GCCGCCCGTC TTGA

214

(75) INFORMATION FOR SEQ ID NO 74:

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- (i) SEQUENCE CHARACTERISTICS:
 - (A) LENGTH: 236
 - (B) TYPE: nucleic acid
 - (C) STRANDEDNESS: single
 - (D) TOPOLOGY: linear
- (ii) MOLECULE TYPE: Trichosporon beigelii specific region of 28S gene.
- (iii) HYPOTHETICAL: No
- (iv) ANTISENSE: No
 - (v) SEQUENCE DESCRIPTION: SEQ ID NO: 74:

AGTCAGTCGT GTTCTTTGGA TTCAGCCAGT TCTGCTGGTC
TACTTCCTTG 50

GAACGGTCA ACATCAGTTT TGTCCGGTGG ATAAAGGTAG
TAGGAATGTG 100

ACTTCTCCGG AAGTGTTATA GCCTATTATC ACATACACTG GGTGAGACTG 150

AGGACTGCAG CTCGCCTTTA TGGCCGGCCT TCGGGCACGT TCGAGCTTAG 200

GATGTTGACA TAATGGCTTT AAACGACCCG TCTTGA

- (76) INFORMATION FOR SEQ ID NO:75:
 - (i) SEQUENCE CHARACTERISTICS:
 - (A) LENGTH: 14
 - (B) TYPE: nucleic acid
 - (C) STRANDEDNESS: single
 - (D) TOPOLOGY: linear

	(ii) WOLLCOLL 1 11 L. Nucleic acid probe for l'aracoccidioides brasiliensis
	(iii) HYPOTHETICAL: No
5	(iv) ANTISENSE: No
	(v) SEQUENCE DESCRIPTION: SEQ ID NO:75:
	ACTCCCCGT GGTC 14
10	(77) INFORMATION FOR SEQ ID NO:76:
	(i) SEQUENCE CHARACTERISTICS:
	(A) LENGTH: 14
15	(B) TYPE: nucleic acid
	(C) STRANDEDNESS: single
	(D) TOPOLOGY: linear
20	(ii) MOLECULE TYPE: Nucleic acid probe for Pneumocystis carinii
20	(iii) HYPOTHETICAL: No
	(iv) ANTISENSE: No
25	(v) SEQUENCE DESCRIPTION: SEQ ID NO:76:
-	GGAAGGGAAA TTGG 14
	(78) INFORMATION FOR SEQ ID NO:77:
30	(i) SEQUENCE CHARACTERISTICS:
	(A) LENGTH: 223

(B) TYPE: nucleic acid

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(C) STRANDEDNESS: single

(D) TOPOLOGY: linear

- (ii) MOLECULE TYPE: Paracoccidioides brasiliensis specific region of 28S gene.
 - (iii) HYPOTHETICAL: No
 - (iv) ANTISENSE: No
 - (v) SEQUENCE DESCRIPTION: SEQ ID NO:77:

GCGCTTGCGA CCAGAGTCGG CCGCGGGGGC TCAGCGGGCA

15 CTCGTTGCCC 50

GTGCACTCCC CCGTGGTCGG GCCAGCGTCG GTTTCGACGG CCGGTCAAAG 100

GCCCCGGAA TGTGTCGCCT CTCGGGGCGT CTTATAGCCG GGGGTGCAAT 150

20 GCGGCCAGTC GGGACCGAGG AACGCGCTCC GGCACGGACG
CTGGCTTAAT 200

GGTCGTAAGC GACCCGTCTT GAA

223

(79) INFORMATION FOR SEQ ID NO:78:

- (i) SEQUENCE CHARACTERISTICS:
 - (A) LENGTH: 257
 - (B) TYPE: nucleic acid
 - (C) STRANDEDNESS: single
 - (D) TOPOLOGY: linear
- (ii) MOLECULE TYPE: Pneumocystis carinii specific region of 28S gene.

(iii) HYPOTHETICAL

- (iv) ANTISENSE: No
- (v) SEQUENCE DESCRIPTION: SEQ ID NO:78:

GCGCTTGTGA TCAGACATGC CTTTATAGGA GATGCCATTG

TTTCGGCATT 50

GGCAGTATTA TCCGAATTGG CAGGCCAGCA TCGGTTTCAG
TTACTGGATA 100

AAACTGGAAG AAGGTAGGCT CTCTTCGGAG GGTTTTTTAG CTTCCAGTAG 150

CTGCAGTGAC CGGGACCGGA AGGGAAATTG GGTCTTTGAA GACCTTATGA 200

TGTTGGCAGA AATGGTCCTA AGCGACCCGT CTTGAAACAC GGACCAAGGA 250

GTCTAAT

257

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(80) INFORMATION FOR SEQ ID NO 79:

- (i) SEQUENCE CHARACTERISTICS:
 - (A) LENGTH: 20
 - (B) TYPE: nucleic acid
 - (C) STRANDEDNESS: single
 - (D) TOPOLOGY: linear
- (ii) MOLECULE TYPE: Nucleic Acid Primer

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(iii) HYPOTHETICAL: No

- (v) SEQUENCE DESCRIPTION: SEQ ID NO: 79:
 - ATCAATAAGC GGAGGAAAAG

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- (81) INFORMATION FOR SEQ ID NO 80:
 - (i) SEQUENCE CHARACTERISTICS:
 - (A) LENGTH: 20
 - (B) TYPE: nucleic acid
 - (C) STRANDEDNESS: single
 - (D) TOPOLOGY: linear

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- (ii) MOLECULE TYPE: Nucleic Acid Primer
- (iii) HYPOTHETICAL: No
- (iv) ANTISENSE: No
 - (v) SEQUENCE DESCRIPTION: SEQ ID NO: 80:

CTCTGGCTTC ACCCTATTC

CLAIMS

We claim:

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- 1. An oligonucleotide hybridization probe for *Paracoccidioides brasiliensis*, said probe having the nucleotide residue sequence of (SEQ ID NO:75) or the complement thereof.
 - 2. An oligonucleotide hybridization probe for *Pneumocystis carinii*, said probe having the nucleotide residue sequence of (SEQ ID NO:76) or the complement thereof.
 - 3. A method of determining whether one or more fungal species selected from the group of fungal species consisting of *Paracoccidioides brasiliensis* and *Pneumocystis* carinii is present in a sample of fungi, said method comprising the following steps:
 - a) extracting nucleic acid material from fungi contained in said sample;
 - b) adding two known oligonucleotide primers, one of said primers being (SEQ ID NO:1) or (SEQ ID NO:2), said primers bracketing a hypervariable region on the 28S rDNA or rRNA present in the fungal species of said group;
 - c) amplifying the sequence between said primers; and
 - d) using one or more labeled probes directed to a portion of the hypervariable region bracketed by said primers, each said labeled probe being specific for one of said fungal species from said group, to determine whether said fungal species identified by each said labeled probe is present in said sample.
 - 4. The method of claim 3 in which, in said amplifying step, said amplifying procedure is the polymerase chain reaction
- 5. The method of claim 3 in which said one or more probes is selected from the group consisting of (SEQ ID NO:75) and (SEQ ID NO:76).

6. The method of claim 3 wherein, in step (d), more than one probe is used, each said probe being connected to (a) a different signal moiety or (b) a moiety which allows separation of said probes.

7. A method of determining whether one or more fungal species selected from the group of fungal species consisting of *Paracoccidioides brasiliensis* and *Pneumocystis* carinii is present in a sample of fungi, said method comprising the following steps:

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- a) extracting nucleic acid material from fungi contained in said sample;
- b) adding a universal fungal probe selected from the group consisting of (SEQ ID NO: 1), (SEQ ID NO:2) and the complements thereof;
- c) using one or more second probes, each said second probe being specific for one of said fungal species from said group, wherein said one or more second probes each has a nucleotide residue sequence selected from the group consisting of (SEQ ID NO:75), (SEQ ID NO:76) and the complements thereof; and
- d) determining whether said fungal species identified by each said second probe is present in said sample, wherein at least one of said probes is connected to a signal moiety and at least one of said probes is connected to a moiety that allows separation of said probes.
- 8. A method of determining whether one or more fungal species selected from the group of fungal species consisting of *Paracoccidioides brasiliensis* and *Pneumocystis carinii* is present in a sample of fungi, said method comprising the following steps:
 - a) extracting nucleic acid material from fungi contained in said sample; and
 - b) using one or more labeled probes, each said labeled probe being specific for one of said fungal species from said group, to determine whether said fungal species identified by each said labeled probe is present in said sample, wherein said one or more labeled probes each has a nucleotide residue sequence selected from the group consisting of (SEQ ID NO:75), (SEQ ID NO:76) and the complements thereof.

9. A species specific reference oligonucleotide for *Paracoccidioides brasiliensis* having the nucleotide residue sequence of SEQ ID NO:77 or the complement thereof.

- 10. A species specific reference oligonucleotide for *Pneumocystis carinii* having the nucleotide residue sequence of SEQ ID NO:78 or the complement thereof.
 - 11. A method of determining whether one or more fungal species selected from the group of fungal species consisting of *Paracoccidioides brasiliensis* and *Pneumocystis carinii* is present in a sample of fungi, said method comprising the following steps:
 - a) extracting nucleic acid material from fungi contained in said sample:
 - b) adding two known oligonucleotide primers, one of said primers being (SEQ ID NO:1) or (SEQ ID NO:2), said primers bracketing a hypervariable region on the 28S rDNA or rRNA present in the fungal species of said group;
 - c) amplifying the sequence between said primers; and
 - d) using one or more labeled probes directed to a portion of the hypervariable region bracketed by said primers, each said labeled probe being specific for one of said fungal species from said group, wherein each said one or more labeled probes is fully complementary to a species-unique nucleotide sequence in said hypervariable region, to determine whether said fungal species identified by each said labeled probe is present in said sample, wherein, furthermore, said one or more labeled probes each has a nucleotide residue sequence consisting of from 10 to 50 consecutive nucleotide residues from a sequence selected from the group consisting of (SEQ ID NO:77) and (SEQ ID NO:78) and the complements thereof.
- 12. A method of claim 11 in which said amplifying procedure is the polymerase chain reaction.

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13. A method of claim 11 wherein more than one third probe is used, each said third probe connected to a different signal moiety or moiety which allows separation of said third probe.

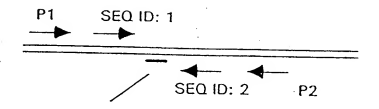
A method of determining whether one or more fungal species selected from the group of fungal species consisting of *Paracoccidioides brasiliensis* and *Pneumocystis carinii* is present in a sample of fungi, said method comprising the following steps:

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- extracting the nucleic acid material from the fungi contained in said sample;
- b) adding a universal fungal probe selected from the group consisting of (SEQ ID NO: 1), (SEQ ID NO:2) and the complements thereof;
- using one or more second probes, each said second probe being specific for one of said fungal species from said group, wherein each said one or more second probes is fully complementary to a species-unique nucleotide sequence in said hypervariable region and wherein, further, said one or more second probes each has a nucleotide residue sequence consisting of from 10 to 50 consecutive nucleotide residues from a sequence selected from the group consisting of (SEQ ID NO:77) and (SEQ ID NO:78) and the complements thereof; and
- d) determining whether said fungal species identified by each said second probe is present in said sample,
 wherein at least one of said probes is connected to a signal moiety and at least one of said probes is connected to a moiety that allows separation of said probes.
 - 15. An oligonucleotide probe/primer for fungi, said probe having the nucleotide residue sequence of (SEQ ID NO:79) or the complement thereof.
 - 16. An oligonucleotide probe/primer for fungi, said probe having the nucleotide residue sequence of (SEQ ID NO:80) or the complement thereof.



SEQ ID: 3 to SEQ ID: 23

FIGURE 1

INTERNATIONAL SEARCH REPORT

Inter nal Application No PCT/IB 98/00865

		PC1	T/IB 98/00865
A. CLASS	FICATION OF SUBJECT MATTER C1201/68		
1,00	01241/00		
A	ha lada ara atau di Bada at Cita a Wasai a Mara		
	to International Patent Classification(IPC) or to both national class	sification and IPC	
Minimum d	ocumentation searched (classification system followed by classifi	cation symbols)	
IPC 6	C12Q		
Documenta	ition searched other than minimum documentation to the extent the	at such documents are included in	the fields searched
Electronic o	data base consulted during the international search (name of dat	a base and, where practical, search	terms used)
	<i>y</i>		
			· ·
C. DOCUM	ENTS CONSIDERED TO BE RELEVANT		
Category *	Citation of document, with indication, where appropriate, of the	relevant passages	Relevant to claim No.
Х	HO OF 21741 A (CIDA CODNING DI	LONGOTIOS	<u>.</u>
^	WO 96 21741 A (CIBA CORNING DIA CORP ;SANDHU GURPREET S (US); I		3,4,6,9, 11-14
	18 July 1996	(LINE ONOUL)	11 14
Y	see the whole document		7,8
Χ	SANDHU G S ET AL: "MOLECULAR I	PROBES FOR	15,16
	DIAGNOSIS OF FUNGAL INFECTIONS		12,21
	JOURNAL OF CLINICAL MICROBIOLOG vol. 33, no. 11, 1 November 199		α
	2913-2919, XP002002934	os, pages	
Y	see the whole document		3,4,6-8,
			11-14
	÷.	-/	
			·
X Furti	her documents are listed in the continuation of box C.	Y Patent family member	s are listed in annex.
° Special ca	tegories of cited documents :	"T" later document published a	after the international filling date
"A" docume consid	ent defining the general state of the art which is not lered to be of particular relevance	or priority date and not in cited to understand the pri invention	conflict with the application but inciple or theory underlying the
"E" earlier o	document but published on or after the international late	"X" document of particular rele	vance; the claimed invention
which	int which may throw doubts on priority claim(s) or is cited to establish the publication date of another	involve an inventive step	vel or cannot be considered to when the document is taken alone
citation	n or other special reason (as specified) ent referring to an oral disclosure, use, exhibition or	"Y" document of particular rele cannot be considered to i document is combined wi	vance; the claimed invention nvolve an inventive step when the th one or more other such docu-
other r "P" docume	neans ant published prior to the international filing date but	ments, such combination in the art.	being obvious to a person skilled
later th	nan the priority date claimed	"&" document member of the s	
Date of the	actual completion of theinternational search	Date of mailing of the inter	national search report
1	September 1998	21/09/1998	
Name and n	nailing address of the ISA	Authorized officer	
	European Patent Office, P.B. 5818 Patentlaan 2 NL - 2280 HV Rijswijk Tel. (+31-70) 340-2040, Tx. 31 651 epo nl, Fax: (+31-70) 340-3016	Knehr, M	

INTERNATIONAL SEARCH REPORT

Inte anal Application No PCT/IB 98/00865

C.(Continuation) DOCUMENTS CONSIDERED TO BE RELEVANT							
C.(Continu Category :		Relevant to claim No.					
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Y	WO 96 29432 A (UNIV BOSTON) 26 September 1996 see the whole document	3,4,6-14					
Υ .	LIU Y ET AL.: "Sequence and variability of the 5.8S and 26S rRNA genes of Pneumocystis carinii" NUCLEIC ACIDS RESEARCH, vol. 20, no. 14, 1992, pages 3763-3772, XP002075979 see abstract see page 3763, column 2, paragraph 2 see page 3766, column 1, paragraph 1 - column 2, paragraph 1 see page 3767, column 2, paragraph 2 see page 3769, column 1, paragraph 2 see page 3770, column 2, paragraph 2 - page 3770, column 2, paragraph 1; figures 1,5,7,8; tables 1,3	3,4,6-14					
ſ	WO 95 29260 A (CIBA GEIGY AG ;LIGON JAMES M (CH); BECK JAMES J (US)) 2 November 1995 see the whole document	3,4,6-8, 11-15					
Y	US 5 501 951 A (MILLIMAN CURT L) 26 March 1996 see abstract see column 2, line 57 - column 3, line 2 see column 3, line 29 - line 51 see column 4, line 21 - line 34 see column 6, line 12 - line 25; table 1	3,4,6-8, 11-15					
A	WO 94 02636 A (HITACHI CHEMICAL CO LTD; MITSUHASHI MASATO (US); COOPER ALLAN (US)) 3 February 1994 see the whole document						
Ρ,Χ	SANDHU G S ET AL.: "Molecular detection and identification of Paracoccidioides brasiliensis" JOURNAL OF CLINICAL MICROBIOLOGY, vol. 35, no. 7, 1997, pages 1894-1896, XP002075980 see the whole document	1,3-8, 11-14					

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Inter and Application No
PCT/IB 98/00865

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